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(71) Applicant: Sasol Technology (Proprietary)
Limited
2196 Johannesburg (ZA)

(72) inventors:

• Joubert, Dawid Johannes
Sasolburg 9570 (ZA)

 Young, Desmond Austin Vanderbijlpark 1911 (ZA)
 Tincul, loan

Sasolburg 9570 (ZA)

(74) Representative: Ablewhite, Alan James MARKS & CLERK, 57/60 Lincoln's Inn Fields London WC2A 3LS (GB)

(54) Polymerization

(57) A polymer of ethylene as a first monomeric component, with a high carbon number linear alpha olefin having at least five carbon atoms as a second monomeric component. At least one of the monomeric components is Fischer- Tropsch derived so that it includes at least one other olefinic component. A process for producing the polymer is also provided.

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Description

- ... [0001] THIS INVENTION relates to polymerization. It relates in particular to a polymer, and to a process for producing a polymer.
- [0002] According to a first aspect of the invention, there is provided a polymer of ethylene as a first monomeric component, with a high carbon number linear alpha olefin having at least five earbon atoms as a second monomeric component, with at least one of the monomeric components being Fischer-Tropsch derived so that it includes at least one other olefinic components.
- [0003] In other words, a polymer according to the first aspect of the invention is the reaction product obtained when a first monomeric component comprising ethylene is reacted with a second monomeric component comprising a high carbon number linear alpha olefin having at least five carbon atoms, with at least one of the monomeric components being Fischer-Tropsch derived so that it includes at least one other olefinic component. Normally a plurality of the other olefinic components will be present in a Fischer-Tropsch derived monomeric component.
- [0004] More particularly, the polymer may be that obtained by reacting the first monomeric component with the second monomeric component in the presence of a metallocene catalyst.
- [0005] The high carbon number linear olefin may be 1-pentene, 1-hexene, 1-heptene, 1-octene or 1-nonene.
- (0006) The first and/or the second monomeric component is thus, as hereinbefore set out, Fischer-Tropsch derived.
 The polymer may include a least one further cledinci monomeric component. Thus, the polymer may include, as a third monomeric component, he light carbon number linear alpha olefin having 4 or more carbon atoms, and which is different to that of the second monomeric component, with the hird monomeric component also being Fischer-Tropsch deviced so that it includes at least one other olefinic component. It may then also, if desired, include at least one further different high carbon number linear alpha olefin which is Fischer-Tropsch derived. In other words, it will then include a pully of Fischer-Tropsch derived monomeric components. Instead, or additionally, however, the polymer may include at least one conventional olefinic monomeric components. Thus, the polymer may comprise at least one conventional olefinic monomeric component to the polymer may include at least one conventional olefinic monomeric component to the polymer with a plurally of Fischer-Tropsch derived in the second monomeric component to register with a plurally of Fischer-Tropsch.
- [0007] By Fischer-Tropsch derived in respect of a monomeric component is meant that it is obtained from the socalled Fischer-Tropsch process, ie it is obtained by reacting a synthesis gas comprising carbon monoxide and hydrogen in the presence of a suitable Fischer-Tropsch catalyst, normally a cobalt, iron, or cobalt/iron Fischer-Tropsch catalyst, at elevated temperature in a suitable reactor, which is normally a fixed or sturry bed reactor, thereby to obtain a range of products, including oldfinic monomeric components suitable for use in the polymers of this invention. The products from the Fischer-Tropsch reaction must then usually be worked up to obtain individual products such as the olefinic monomeric components suitable for use in the polymers of the present invention.
- [0008] By 'conventional olefinic monomeric component' is meant any olefinic monomer that is not Fischer Tropsch derived and that can be used in copolymenzation with low carbon number olefins.
- [0009] Thus, the polymers according to this invention may be polymers of ethylene, with at least one Fischer Tropsch derived linear alpha olelin or with a mixture of a linear alpha olelin obtained from a Fischer-Tropsch process and any other polymerization grade olelinic monomer(s) obtained from other processes, provided that these polymers are obtained by polymerization of the olelins or monomer(s) in the presence of a metallocene catalyst, or are the product of reaction of the olelins or monomer(s) in the presence of a metallocene catalyst.
- [0010] The inventors surprisingly discovered that when the olefinic monomers employed in catalyzed polymerization as the second monomeric component, are obtained from the Fisher-Tropsch process, the resultant polymers have very large domains of fundamental and/or application properties, and may be superior in some of these properties to those of polymers in which all the monomers have been obtained by conventional methods. The inventors believe that this unexpected behavior is due to very small amounts of the other olefinic components present in the Fischer Tropsch derived olefinic component and which until now have been regarded as impurities. These other olefinic components may be other hydrocarbons having one or more double bonds, whether linear, branched or aromatic, with the exception of those which poison the catalyst to the extent that it no longer polymerizes the monomers. The inventors further believe that these components may sendent as a continuous control of the service of the component and the polymers obtained according to this invention, thus improving the processability of these polymers. These components may selectively and/or partially and/or temporarily modify the initiation of ethylene polymerization or the insertion of the ethylene in the growing chain or the termination of polymertization, thereby changing the distribution of the components in the polymer chain and/or the content level of the individual components in the polymer and/or the circuit distribution, and/or the moticular weight of the polymer chain and/or the content level of the individual components in the polymer and/or the english of branching of the polymer and/or the moticular depolyment and/or the moticular weight of the polymer and/or the moticular depolyment.
- sequence length and/or its morphology, with any one or more of these being reflected in unexpected application properties of the resultant polymers.

 [0011] However, the inventors have also discovered that, for practical applications when the linear alpha oldfinic monomers employed in the polymerization as the second-monomeric component, are obtained from the Fisher-Tropseh

process, the proportion of the other olefinic components referred to hereinbefore in the second monomer component is preferably within particular limits.

- [0012] Thus, the amount of these other oldfinic components present in the second monomeric component, when obtained from the Fisher-Tropsch process, may be from 0,002% to 2%, more preferably from 0,02% to 2%, and most preferably from 0,2% to 2%, based on the total mass of the monomeric component, le given on a mass or weight basis. However it is to be noted that in particular cases the total amount of the other oleflinic components in the monomeric component may be above the limits hereinbefore set out.
- [0013] The ethylene may also be obtained from the Fischer-Tropsch process. However, due to the process of separation and purification involved in obtaining Fischer-Tropsch derived ethylene, polymers containing Fischer-Tropsch of derived ethylene may, in certain cases, not show any difference to polymers containing ethylene obtained from conventional processes.
 - [0014] The second monomeric component or Fischer Tropsch derived comonomer may be a linear alpha olefin having a total number of carbon atoms between 5 and 9, leading thus to different groups of polymers. Typical examples of such high carbon number linear alpha olefins are 1-pentieng, 1-hexene, 1-heptene, 1-checene and 1-nonement.
- 15 [0015] When the high carbon number linear alpha olefin is 1-pentene, the other olefinic impurities typically comprise mainly:
 - 2-methyl-1-butene up to 0.46%; and/or

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- branched olefins having a carbon number of 5; and/or
- Internal olefins having a carbon number of 5: and/or
 - cyclic olefins having a carbon number of 5.

[0016] When the high carbon number linear alpha clefin is 1-hexene, the other clefinic impurities typically comprise mainly:

- branched olefins, mainly having a carbon number of 6 up to 0,51%; and/or
- Internal olefins, mainly having a carbon number of 6 up to 0,18%; and/or
- cyclic olefins, mainly having a carbon number of 6 up to 0,13%.
- 30 [0017] When the high carbon number linear olefin is 1-heptene, the other olefinic impurities typically comprise mainly:
 - branched olefins, mainly having a carbon number of 7 up to 0,48%; and/or
 - Internal olefins, mainly having a carbon number of 7 up to 0,53%.
- 35 [0018] When the high carbon number linear alpha olefin is 1-octane, the other olefinic impurities typically comprise mainly:
 - branched olefins, mainly having a carbon number of 8 up to 0,41%; and/or
 - Internal olefins, mainly having a carbon number of 8 up to 0,83%.
 - [0019] When the high carbon number linear alpha olefin is 1-nonene, the other olefinic impurities typically comprise mainly:
- branched olefins, mainly having a carbon number of 9 up to 0,65%; and/or
- internal olefins, mainly having a carbon number of 9 up to 0,51%.

[0020] More specifically, according to the first aspect of the invention, there is thus provided a polymer of ethylene as the first monomeric component, with a high carbon number linear alpha leight having at least two earthon atoms as the second-monomeric component and which is obtained from a Flischer-Tropsch process or is Flischer-Tropsch derived, with this polymer being obtained by copolymerization in the presence of a metallicone catalyst or being the recipion product of ethylene and the second monomeric component which is obtained from a Fischer-Tropsch process or is Fischer-Tropsch derived, in the presence of a metallicone catalyst.

[0021] The ratio of the molar proportion of the ethylene to the molar proportion of the component votice is obtained from a Fischer Tropsch process or is Fischer Tropsch derived may be from 99.9:

55 0, 1 to 90:20. The preferred ratio of the molar proportion is from 99.9:0,1 to 90:10. The most preferred ratio of the molar proportion is from 99.9:0,1 to 90:10. The most preferred ratio of the molar proportion is from 99.9:0.1 to 90:10. The most preferred ratio of the molar proportion is from 99.9:0,1 to 90:10. The most preferred ratio of the molar proportion is from 99.9:0.1 to 96:5.

[0022] Typical examples of Fischer-Tropsch derived olefins which can be used in the different embodiments of the invention are those as hereinbefore described in respect of this aspect of the invention, and which typically have levels

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of other olefinic components present therein as hereinbefore described.

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[0023] Thus, in one embodiment of this aspect of the invention, the component or Fischer-Tropsch derived second monomenic component may comprise from 0,002% to 2%, by mass, other olefinic components.

[0024] In another embodiment of this aspect of the invention, it may comprise from 0,02% to 2%, by mass, other olefinic components.

[0025] In yet another embodiment of this aspect of the invention, it may comprise from 0,2% to 2%, by mass, other olefinic components

[0026] In a still further embodiment of this aspect of the invention, it may comprise from 0,2% to in excess of 2%, by

mass, other olefinic components.

[0027] In particular, the polymer may be that obtained by reacting at least ethylene, with the comonomer or second monomeric component which is obtained from a Fischer Tropsch process or is Fischer Tropsch derived in one or more reaction zones, while maintaining in the reaction zone(s) a pressure in the range between atmospheric pressure and 5000kg/cm2 and a temperature between ambient and 300°C, in the presence of a catalyst or catalyst system comprising a catalyst and a cocatalyst. The catalyst may in particular, be a metallocene catalyst.

[0028] The inventors have surprisingly found that in the families of copolymers of ethylene with a high carbon number Fischer-Tropsch derived alpha olefin or second monomeric component which is obtained from a Fischer-Tropsch process or is Fischer-Tropsch derived, there can be found particular distinguishable subfamilies of polymers with a large range of unexpected properties dependent on the different second comonomers or olefinic components which are Fischer-Tropsch derived or obtained from a Fischer-Tropsch process, having different numbers of total carbon atoms,

hoen [0029] The properties of the polymers in each family and subfamily group are determined mainly by the ratio of the proportion of ethylene to that of the second monomeric component which Fischer-Tropsch derived or obtained from a Fischer Tropsch process, and the amounts of the other olefinic components contained in the second monomeric component. In this manner, a large range of particular polymers can be obtained, having a large range of application properties controlled between certain limits. The resultant polymers are suitable for improved application in the main

processing fields. Typical applications of the terpolymer include extrusions, blow moulding and injection moulding. [0030] The polymer according to the first aspect of the invention for each embodiment hereinbefore described may have the following properties:

20 a) a melt flow rate as measured according to ASTM D 1238 in the range of 0.01 to about 100g/10min; and/or b) a density measured according to ASTM D 1505 in the range of about 0.835 to about 0.950.

[0031] In a first version of each embodiment of this aspect of the invention, the polymer may be that obtained by the reaction of ethylene with 1-pentene

[0032] In a second version of each embodiment of this aspect of the invention, the polymer may be that obtained by the reaction of ethylene with 1-hexene [0033] In a third version of each embodiment of this aspect of the invention, the polymer may be that obtained by

the reaction of ethylene with 1-heptene [0034] In a fourth version of each embodiment of this aspect of the invention, the polymer may be that obtained by

the reaction of ethylene with 1-octene [0035] In a fifth version of each embodiment of this aspect of the invention, the polymer may be that obtained by the

reaction of ethylene with 1-nonene. [0036] The inventors surprisingly discovered that the copolymers according to this invention are characterized by

particular rheological properties.

[0037] Rheological analysis of the polymers were performed on a Physica MCR-500 rheometer with a parallel plate measuring system (25 mm diameter, 1 mm gap), under controlled strain conditions (strain amplitude 1%) at a constant temperature for three particular temperatures (150°C, 160°C, 170°C). The Carreau-Gahleitner parameters of dynamic zero shear viscosity η°, dynamic Infinite shear viscosity η_{Inf}, characteristic relaxation time a and power exponents b, p and n were obtained using the rheology software.

[0038] The polymer may thus have a power exponent b which complies with the following equations, namely:

At 150°C, b ≥ 0.0437(C) + 0.2013

At 160 °C, b ≥ 0.0308[C] + 0.2138

At 170°C, b ≥ 0.0308[C] + 0.2538,

where b is a Carreau-Gahleitner parameter, and [C] is the high carbon number linear alpha olefin content in mole 5 percentage.

[0039] The polymer may have a power exponent p which complies with the following equations, namely:

At 150°C, p ≥ -1.2877[C] + 6.8666

At 160°C, p ≥ -1.1233[C] + 6.3942

At 170°C, p ≥ -1.1507(C) + 6.3063.

where p is a Carreau-Gahleitner parameter, and [C] is the high carbon number linear alpha olefin content in mole percentage.

The polymer may have a power exponent n which complies with the following equations, namely:

At 150°C, n ≥ 0.2995[C] - 0.8328

At 160°C, n ≥ 0.3011[C] - 0.8435

At 170°C, n ≥ 0.2942[C] - 0.8115

where n is a Carreau-Gahleitner parameter, and [C] is the high carbon number linear alpha olefin content in mole percentage.

[0040] The polymer may also comply with the following equations, namely:

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At 150°C, 1/MFI ≥ 1.5364e-1E-06η°

At 160°C, 1/MFI ≥ 1.5486e-1E-08η*

At 170°C, 1/MFI ≥ 1.5513e^{-1E-06}η°

where MFI is the melt flow index and η° is the dynamic zero shear viscosity.

[0041] According to a second aspect of the invention, there is provided a process for producing a polymer, which comprises reacting at least a first monomeric component comprising a bright carbon number linear alpha olefin having at least two carbon atoms, and wherein at least one of the monomeric components is Fischer-Tropsch derived so that it contains also one or more other olefinic components, in one or more reaction zones, while maintaining the reaction zones, whose between atmospheric pressure and 5000kg/cm², and at a temperature between ambient and 300°C, in the presence of a catalyst, or a catalyst system comprising a catalyst and so cata

[0042] In a first embodiment of this aspect of the invention, the high carbon number linear alpha olefin may be added at the start of the reaction while the ethylene is added continuously during the course of the reaction.

[0043] In a second embodiment of this aspect of the invention, the reaction may be effected in a continuous fashion, with the elthylene being added continuously, and with the high carbon number linear alpha olefin being added continuously during the course of the reaction.

[0044] In a third embodiment of this aspect of the invention, the reaction may be effected in a continuous fashion, with the othylene being added continuously, and with the high carbon number linear alpha olefin being added discontinuously during the course of the reaction.

- [0045] In a fourth embodiment of this aspect of the invention, the reaction may be effected in a continuous fashion, with the ethylene being added discontinuously, and with the high carbon number linear alpha clefin being added continuously during the course of the reaction.
- [0048] According to a third aspect of the invention, there is provided a polymer of ethylene as a first monomeric component, with a first high carbon number linear alpha olefin having at least five earbon a lotnes as a second monomeric component and with a second different high carbon number linear alpha olefin having at least four carbon atoms as a third monomeric component, with at least one of the monomeric components being Fischer-Tropsch darivad so that it contains also at least one other olefinic component.
- [0047] In other words, according to the third aspect of the Invention, there is provided a polymer which is the reaction product of ethylene as a first monometric component with a first high extenon number linear alpha olefth having at least five action atoms as a second monometric component and at least one different high earbon number linear alpha leafth having at least four carbon atoms as a third monometric component, with at least one of the monometric components being Fischar-Tropsch derived.
- [0048] Yet further, according to the third aspect of the Invention, there is thus provided a terpolymer of othylene as a first monomeric component with a first high carbon number linear alpha olefin having at least five carbon atoms as a second monomeric component and at least on different high carbon number linear alpha olefin having at least five carbon atoms as a third monomeric component, with at least one of the monomeric components being Fischer-Tropsch derived.
- [0049] Still further, according to this aspect of the invention, there is provided a polymer of ethylene with at least two different high carbon number linear alpha olefins each having at least four carbon atoms.
- [0050] The ratio of the motar proportion of the ethylene to the sum of the motar proportions of the high carbon number linear alpha oldfism may be from 99,90,1 to 80:20. The preferred ratio of the motar proportions is from 99,9:0,1 to 90:10. The most preferred ratio of the motar proportions for my 99,9:0,1 to 95:5.
- [0051] Typical examples of Fischer-Tropsch derived high carbon number olefins which can be used in the different ambodiments of the invention are those as harainbefore described in respect of the first aspect of the invention, and which typically have levels of other olefinic components present therein as hereinbeford described.
 - which typically have levels or other olerinic components present therein as hereinbefora described.

 [0052] Thus, In one embodiment of this aspect of the invention, each Fischer-Tropsch derived monomeric component may comprise from 0.002% to 2% by mass, other olefinic components.
- [0053] In another embodiment of this aspect of the invention, it may comprise from 0,02% to 2%, by mass, other olefinic components.
 - [0054] In yet another embodiment of this aspect of the invention, it may comprise from 0,2% to 2%, by mass, other olefinic components.
 - [0055] In a still further embodiment of this aspect of the invention, it may comprise from 0,2% to in excess of 2%, by mass, other olefinic components.
 - [0056] In particular, the polymer may be that obtained by reacting ethylene, with two or more high carbon number linear alpha olders or components of which at least one is Fischer-Tropseh derived, in one or more reaction zones, while maintaining in the reaction zone(s) a pressure in the range between atmospheric pressure and 5000kg/cm² and a temporatrue between armisent and 300°C, in the presence of a metallecence catalyst.
- [0057] The inventors have surprisingly found that in the family of tarpolymers of athylena with two or mora high or carbon number linear alpha olefins or components of which at least one is Fischer-Tropsch derived, there can be found particular distinguishable groups of polymers with a large range of unexpected properties depondent on the different Fischer-Tropsch derived components or monomeric components, having different numbers of total carbon atoms,
- [0058] The properties of the tempolymers in each family and subfamily group are determined mainly by the ratio of the proportion of einhylene to the sum of the commonenes or Fischer-Tropsch derived monomeric components, and the amounts of the other olderlinic components contained in the Fischer-Tropsch derived monomeric components.
- [0059] However the properties are also detarmined by the ratio of molar proportions of the different high carbon number linear alpha olefins of which at least one is Fischer-Tropsch derived.
- [0060] The ratio of the molar proportions of the different high carbon number linear alpha clefins of which at least one is Fischer-Tropsch derived may be from 0,1:99,9 to 99,9:0.1, more preferably from 1:99 to 99:2, and more preferably from 2:98 to 98:2.
 - [0081] In this manner, an even larger range of particular polymers can be obtained than in the first aspect of the invention, having a large range of application properties controlled between certain limits. The relating the suitable for improved application in the main processing fields. Typical applications of the terpolymer include extrusions, blow moudifies and inlection moulding.
 - [0062] The polymer according to the third aspect of the invention for each embodiment hereinbefore described may have the following properties:

- a) a melt flow rate as measured according to ASTM D 1238 in the range of 0,01 to about 100g/10min; and/or
 b) a density as measured according to ASTM D 1505 in the range of about 0,835 to about 0,950.
- [0063] In a first version of each embodiment of this aspect of the invention, the polymer may be that obtained by the reaction of ethylene with 1-pentene and a third different alpha olefin.
 - [0064] In a second version of each embodiment of this aspect of the invention, the polymer may be that obtained by the reaction of ethylene with 1-hexene and a third different alpha olefin.
- [0065] In a third version of each embodiment of this aspect of the invention, the polymer may be that obtained by the reaction of ethylene with 1-heptene and a third different alpha olefin.
- [0066] In a fourth version of each embodiment of this aspect of the invention, the polymer may be that obtained by the reaction of ethylene with 1-octene and a third different alpha olefin.
 - [0067] In a fifth version of each embodiment of this aspect of the invention, the polymer may be that obtained by the reaction of ethylene with 1-nonene and a third different alpha olefin.
- [0068] The inventors surprisingly discovered that the terpolymers according to this aspect of the invention are characterized by particular rheological properties.
- [0069] The terpolymer may have a power exponent b which complies with the following equations:

At 150°C, b > 0.037[C] + 0.2052

At 160°C, b ≥ 0.0395fCl + 0.2342

At 170°C, b ≥ 0.0494[C] + 0.2202,

where b is a Carreau-Gahleitner parameter, and [C] is the high carbon number linear alpha olefin content in mole percentage.

[0070] The terpolymer may have a power exponent p which complies with the following equations:

At 150°C, p ≥ -0.4075[C] + 3.4135

At 160°C, p \geq -0.5016[C] + 3.732

At 170°C, p ≥ -0.9091[C] + 5.3455

40 where p is a Carreau-Gahleitner parameter, and [C] is the high carbon number linear alpha olefin content in mole percentage.

[0071] The terpolymer may have a power exponent n which compiles with the following equations:

At 150°C, n ≥ 0.1 69[C] - 0.3439

At 160°C, n ≥ 0.1765[C] - 0.4004

At 170°C, n ≥ 0.242[C] - 0.6489

where n is a Carreau-Gahleitner parameter, and [C] is the high carbon number linear alpha olefin content in mole percentage.

[0072] The terpolymer may comply with the following equations:

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At 150°C, 1/MFI ≥ 0.123e^{2E-05η°}

At 160°C, 1/MFI ≥ 0.12390^{2E-05}η°

At 170°C, 1/MFI ≥ 0.1275e^{2E-05}η*

where MFI is the melt flow index and no is the dynamic zero shear viscosity.

[0073] The terpolymer according to this aspect of the invention, may be that that obtained by reacting at least ethylene, the first high carbon number linear alpha olefin in one or more reaction zones, while maintaining the reaction zone(s) at a pressure between atmospheric pressure and 5000kg/cm², and at a temperature batwaen amblent and 300°C, in the presence of a catalyst, or a catalyst system comprising a catalyst and coctativest.

[0074] The catalyst may, in particular, be a matallocene catalyst.

[0075] According to a fourth aspect of the Invention, there is provided a process for producing a polymer, which comprises reaching at least ethylene as a first monometic component, a first high carbon number linear alpha olefin having at least five carbon atoms as a second monometic component and a different high carbon number linear alpha olefin having at least flour carbon atoms as third monometic component, with at least one of the monometic component, with at least one of the monometic components being Fischer-Tropsch derived so that it contains also one or more checific components, in one or more reaction zones, while maintaining the reaction zone(s) at a pressure between atmospheric pressure and 500kg/cm², and at a temperatura between ambient and 500°C, in tha presence of a catalyst, or a catalyst system comprising a catalyst and acceptable.

[0076] The catalyst may, in particular, be a metallocene catalyst.

[0077] In a first embodiment of this aspect of the invention, the high carbon number linear alpha olefins may be added simultaneously at the start of the reaction, while the ethylane is added continuously during the course of the reaction.

[0078] In a second embodiment of this aspect of the invantion, one of the high carbon numbar linear siphs orlents may be added at the start of the reaction while theytene is added continuously during the reaction, with a continuous or discontinuous supply of the other high carbon number linear siphs orlent being provided, and with no product being removed during the reaction.

[0079] In a third embodiment of this aspect of the invention, the reaction may be effected in a continuous fashion, with the ethylene being added continuously, and with the high carbon number linear alpha olefins being added together and continuously during the course of the reaction.

[0080] In a fourth embodiment of this aspect of the invention, the reaction may be effected in a continuous fashion, with the shylene being added continuously, and with the high carbon number linear alpha clefins being added separately and continuously during the course of the reaction.

[0081] In a fifth embodiment of this aspect of the Invention, the reaction may be effected in a continuous fashion, with the ethylene being added continuously, and with the high carbon number linear alpha olefins baing added together but discontinuously during the course of the reaction.

[0082] In a sixth embodiment of this aspect of the invention, the reaction is affected in a continuous fashion, with tha ethylene being added continuously, and with the high carbon number linear alpha olefins being added separately and discontinuously during the course of the reaction.

[0083] Any suitable metallocene catalyst for ethylene polymarization can, at least in principle, be used. Examples of metallocenes which can be used are Group IV transition metallocenes (titanocenes, zircnocenes, harfocenes), which are characterized by two bulky cyclopentadienyl (Op) or substituted cyclopentadienyl ligands (Op) were the substituent may be linear or branched alkyl groups, substituted and run-substituted aromatic and cyclic alighatic groups, metal-locenes with two Op ligands arranged in a chiral array and which may be connected together with chemical bonds by a bridging group. The bridging group may be a linear, branched or aromatic or alighatic carbon containing from 1 to 50 carbon atoms, germanium or silyl groups substituted with linear or branched alkyl groups, substituted and unsubstituted aromatic and cyclic alighatic groups. Table 1 shows a non-limiting list of metallocenes which can in principle be used.

Table 1.

List of Metallocenes		
Cp ₂ ZrCl ₂ (n-BuCp) ₂ ZrCl ₂ (t-BuCp) ₂ ZrCl ₂	(Me ₂ Cp) ₂ ZrMe ₂ (Me ₅ Cp) ₂ ZrMe ₂ Cp ₂ ZrClMe	

Table 1. (continued)

	continued)
List of Metallocenes	
(i-BuCp) ₂ ZrCl ₂	(n-BuCp) ₂ ZrClMe
(n-Bu ₂ Cp) ₂ ZrCl ₂	(t-BuCp) ₂ ZrClMe
(t-Bu ₂ Cp) ₂ ZrCl ₂	(i-BuCp) ₂ ZrClMe
(i-Bu ₂ Cp) ₂ ZrCl ₂	(n-Bu ₂ Cp) ₂ ZrClMe
(n-Bu ₅ Cp) ₂ ZrCl ₂	(t-Bu ₂ Cp) ₂ ZrClMe
(t-Bu ₅ Cp) ₂ ZrCl ₂	(I-Bu ₂ Cp) ₂ ZrCIMe
(I-Bu ₅ Cp) ₂ ZrCl ₂	(n-Bu ₅ Cp) ₂ ZrClMe
(n-PrCp) ₂ ZrCl ₂	(t-Bu ₅ Cp) ₂ ZrClMe
(t-PrCp) ₂ ZrCl ₂	(I-Bu ₅ Cp) ₂ ZrCIMe
(I-PrCp) ₂ ZrCl ₂	(n-PrCp) ₂ ZrClMe
(n-Pr ₂ Cp) ₂ ZrCl ₂	(t-PrCp) ₂ ZrClMe
(t-Pr ₂ Cp) ₂ ZrCl ₂	(i-PrCp) ₂ ZrClMe
(i-Pr ₂ Cp) ₂ ZrCl ₂	(n-Pr ₂ Cp) ₂ ZrClMe
(n-Pr ₅ Cp) ₂ ZrCl ₂	(t-Pr ₂ Cp) ₂ ZrCiMe
(t-Pr ₅ Cp) ₂ ZrCl ₂	(i-Pr ₂ Cp) ₂ ZrClMe
(I-Pr ₅ Cp) ₂ ZrCl ₂	(n-Pr ₅ Cp) ₂ ZrClMe
(PhCp) ₂ ZrCl ₂	(t-Pr ₅ Cp) ₂ ZrClMe
(Ph ₂ Cp) ₂ ZrCl ₂	(I-Pr ₅ Cp) ₂ ZrClMe
(MeCp) ₂ ZrCl ₂	(PhCp) ₂ ZrClMe
(Me ₂ Cp) ₂ ZrCl ₂	(Ph ₂ Cp) ₂ ZrClMe
(Me ₅ Cp) ₂ ZrCl ₂	(MeCp) ₂ ZrClMe
Cp ₂ ZrMe ₂	(Me ₂ Cp) ₂ ZrCIMe
(n-BuCp) ₂ ZrMe ₂	(Me ₅ Cp) ₂ ZrClMe
(t-BuCp) ₂ ZrMe ₂	Cp ₂ TiCl ₂
(i-BuCp) ₂ ZrMe ₂	(n-BuCp) ₂ TiCl ₂
(n-Bu ₂ Cp) ₂ ZrMe ₂	(t-BuCp) ₂ TiCl ₂
(t-Bu ₂ Cp) ₂ ZrMe ₂	(i-BuCp) ₂ TiCl ₂
(i-Bu ₂ Cp) ₂ ZrMe ₂	(n-Bu ₂ Cp) ₂ TiCl ₂
(n-Bu ₅ Cp) ₂ ZrMe ₂	(t-Bu ₂ Cp) ₂ TiCl ₂
(t-Bu ₅ Cp) ₂ ZrMe ₂	(i-Bu ₂ Cp) ₂ TiCl ₂
(I-Bu ₅ Cp) ₂ ZrMe ₂	(n-Bu ₅ Cp) ₂ TiCl ₂
(n-PrCp) ₂ ZrMe ₂	(t-Bu _B Cp) ₂ TiCl ₂
(t-PrCp) ₂ ZrMe ₂	(I-Bu ₅ Cp) ₂ TiCl ₂
(i-PrCp) ₂ ZrMe ₂	(n-PrCp) ₂ TiCl ₂
(n-Pr ₂ Cp) ₂ ZrMe ₂	(t-PrCp) ₂ TiCl ₂
(t-Pr ₂ Cp) ₂ ZrMe ₂	(i-PrCp) ₂ TiCl ₂
(i-Pr ₂ Cp) ₂ ZrMe ₂	(n-Pr ₂ Cp) ₂ TiCl ₂
(n-Pr ₅ Cp) ₂ ZrMe ₂	(t-Pr ₂ Cp) ₂ TICl ₂
(t-Pr ₅ Cp) ₂ ZrMe ₂	(i-Pr ₂ Cp) ₂ TiCl ₂
(i-Pr ₅ Cp) ₂ ZrMe ₂	(n-Pr ₅ Cp) ₂ TiCl ₂
(PhCp) ₂ ZrMe ₂	(t-Pr ₅ Cp) ₂ TiCl ₂
(Ph ₂ Cp) ₂ ZrMe ₂	(I-Pr ₅ Cp) ₂ TiCl ₂
(MeCp) ₂ ZrMe ₂	(PhCp) ₂ TiCl ₂
(Ph ₂ Cp) ₂ TiCl ₂	(t-Pr ₅ Cp) ₂ TiCiMe
(MeCp) ₂ TiCl ₂	(I-Pr ₅ Cp) ₂ TICIMe
(Me ₂ Cp) ₂ TiCl ₂	(PhCp) ₂ TiClMe
(Me ₅ Cp) ₂ TiCl ₂	(Ph ₂ Cp) ₂ TiClMe
Cp ₂ TiMe ₂	(MeCp) ₂ TiClMe
(n-BuCp) ₂ TiMe ₂	(Me ₂ Cp) ₂ TiCiMe
(t-BuCp) ₂ TiMe ₂	(Me ₅ Cp) ₂ TiClMe

Table 1. (continued)

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lable 1.	(continued)
List of Metallocenes	
(i-BuCp) ₂ TiMe ₂	Cp ₂ HfCl ₂
(n-Bu ₂ Cp) ₂ TiMe ₂	(n-BuCp) ₂ HfCl ₂
(t-Bu ₂ Cp) ₂ TiMe ₂	(t-BuCp) ₂ HfCl ₂
(i-Bu ₂ Cp) ₂ TiMe ₂	(I-BuCp) ₂ HfCl ₂
(n-Bu ₅ Cp) ₂ TiMe ₂	(n-Bu ₂ Cp) ₂ HfCl ₂
(t-Bu ₅ Cp) ₂ TiMe ₂	(t-Bu ₂ Cp) ₂ HfCl ₂
(I-Bu ₅ Cp) ₂ TiMe ₂	(i-Bu ₂ Cp) ₂ HfCl ₂
(n-PrCp) ₂ TiMe ₂	(n-Bu ₅ Cp) ₂ HfCl ₂
(t-PrCp) ₂ TiMe ₂	(t-Bu ₅ Cp) ₂ HfCl ₂
(I-PrCp) ₂ TIMe ₂	(I-Bu ₅ Cp) ₂ HfCl ₂
(n-Pr ₂ Cp) ₂ TiMe ₂	(n-PrCp) ₂ HfCl ₂
(t-Pr ₂ Cp) ₂ TiMe ₂	(t-PrCp) ₂ HfCl ₂
(i-Pr ₂ Cp) ₂ TiMe ₂	(i-PrCp) ₂ HfCl ₂
(n-Pr ₅ Cp) ₂ TiMe ₂	(n-Pr ₂ Cp) ₂ HfCl ₂
(t-Pr ₅ Cp) ₂ TiMe ₂	(t-Pr ₂ Cp) ₂ HfCl ₂
(i-Pr ₅ Cp) ₂ TiMe ₂	(i-Pr ₂ Cp) ₂ HfCl ₂
(PhCp) ₂ TiMe ₂	(n-Pr ₅ Cp) ₂ HfCl ₂
(Ph ₂ Cp) ₂ TiMe ₂	(t-Pr ₅ Cp) ₂ HfCl ₂
(MeCp) ₂ TiMe ₂	(I-Pr ₅ Cp) ₂ HfCl ₂
(Me ₂ Cp) ₂ TiMe ₂	(PhCp) ₂ HfCl ₂
(Me ₅ Cp) ₂ TiMe ₂	(Ph ₂ Cp) ₂ HfCl ₂
Cp ₂ TiCiMe	(MeCp) _p HfCl ₂
(n-BuCp) ₂ TiClMe	(Me ₂ Cp) ₂ HfCl ₂
(t-BuCp) ₂ TiClMe	(Me ₅ Cp) ₂ HfCl ₂
(i-BuCp) ₂ TiCiMe	Cp ₂ HfMe ₂
(n-Bu ₂ Cp) ₂ TiClMe	(n-BuCp) ₂ HfMe ₂
(t-Bu ₂ Cp) ₂ TiClMe	(t-BuCp) ₂ HfMe ₂
(i-Bu ₂ Cp) ₂ TiClMe	(i-BuCp) ₂ HfMe ₂
(n-Bu ₅ Cp) ₂ TiClMe	(n-Bu ₂ Cp) ₂ HfMe ₂
(t-Bu ₅ Cp) ₂ TiClMe	(t-Bu ₂ Cp) ₂ HfMe ₂
(i-Bu ₅ Cp) ₂ TiClMe	(I-Bu ₂ Cp) ₂ HfMe ₂
(n-PrCp) ₂ TICIMe	(n-Bu ₅ Cp) ₂ HfMe ₂
(t-PrCp) ₂ TICIMe	(t-Bu ₅ Cp) ₂ HfMe ₂
(i-PrCp) ₂ TiClMe	(i-Bu ₅ Cp) ₂ HfMe ₂
(n-Pr ₂ Cp) ₂ TiClMe	(n-PrCp) ₂ HfMe ₂
(t-Pr ₂ Cp) ₂ TiClMe	(t-PrCp) ₂ HfMe ₂
(i-Pr ₂ Cp) ₂ TICIMe	(I-PrCp) ₂ HfMe ₂
(n-Pr ₅ Cp) ₂ TiCIMe	(n-Pr ₂ Cp) ₂ HfMe ₂
(t-Pr ₂ Cp) ₂ HfMe ₂	[O(SIMe ₂ Cp) ₂]TICl ₂
(i-Pr ₂ Cp) ₂ HfMe ₂	[O(SiMe ₂ t-BuCp) ₂]TiCl ₂
(n-Pr ₅ Cp) ₂ HfMe ₂	Ind ₂ HfCl ₂
(t-Pr ₅ Cp) ₂ HfMe ₂	(2-MeInd) ₂ HfCl ₂
(I-Pr ₅ Cp) ₂ HfMe ₂	(neomenthylCp) ₂ HfCl ₂
(PhCp) ₂ HfMe ₂	(C ₅ Me ₄ Et) ₂ HfCl ₂
(Ph ₂ Cp) ₂ HfMe ₂	[O(SIMe ₂ Cp) ₂]HfCl ₂
(MeCp) ₂ HfMe ₂	[O(SiMe, t-BuCp),]HfCl,
(Me ₂ Cp) ₂ HfMe ₂	Ind ₂ ZrCiMe
(Me ₅ Cp) ₂ HfMe ₂	(2-MeInd) ₂ ZrCIMe
Cp ₂ HfClMe	(neomenthylCp) ₂ ZrClMe
(n-BuCp) ₂ HfClMe	(C ₅ Me ₄ Et) ₂ ZrClMe

Table 1. (continued)

Table 1.	(continued)
List of Metallocenes	
(t-BuCp) ₂ HfCIMe	[O(SiMe ₂ Cp) ₂]ZrClMe
(i-BuCp) ₂ HfClMe	[O(SiMe ₂ t-BuCp) ₂]ZrClMe
(n-Bu ₂ Cp) ₂ HfClMe	Ind ₂ TiClMe
(t-Bu ₂ Cp) ₂ HfClMe	(2-MeInd) ₂ TiCIMe
(i-Bu ₂ Cp) ₂ HfClMe	(neomenthylCp) ₂ TiClMe
(n-Bu ₅ Cp) ₂ HfClMe	(C ₅ Me ₄ Et) ₂ TiClMe
(t-Bu ₅ Cp) ₂ HfClMe	[O(SiMe ₂ Cp) ₂]TiClMe
(i-Bu ₅ Cρ) ₂ HfClMe	[O(SiMe ₂ t-BuCp) ₂]TiClMe
(n-PrCp) ₂ HfClMe	Ind ₂ HfCiMe
(t-PrCp) ₂ HfCIMe	(2-MeInd) ₂ HfCIMe
(i-PrCp) ₂ HfClMe	(neomenthylCp) ₂ HfClMe
(n-Pr ₂ Cp) ₂ HfClMe	(C ₅ Me ₄ Et) ₂ HfClMe
(t-Pr ₂ Cp) ₂ HfClMe	[O(SiMe ₂ Cp) ₂]HfClMe
(i-Pr ₂ Cp) ₂ HfClMe	[O(SiMe ₂ t-BuCp) ₂]HfClMe
(n-Pr ₅ Cp) ₂ HfClMe	Ind ₂ ZrMe ₂
(t-Pr ₅ Cp) ₂ HfClMe	(2-MeInd) ₂ ZrMe ₂
(i-Pr ₅ Cp) ₂ HfCIMe	(neomenthylCp) ₂ ZrMe ₂
(PhCp) ₂ HfClMe	(C ₅ Me ₄ Et) ₂ ZrMe ₂
(Ph ₂ Cp) ₂ HfClMe	[O(SiMe ₂ Cp) ₂]ZrMe ₂
(MeCp) ₂ HfClMe	[O(SiMe ₂ t-BuCp) ₂]ZrMe ₂
(Me ₂ Cp) ₂ HfCIMe	Ind ₂ TiMe ₂
(Me _s Cp) ₂ HfClMe	(2-MeInd) ₂ TiMe ₂
Ind ₂ ZrCl ₂	(neomenthylCp) ₂ TiMe ₂
(2-Melnd) ₂ ZrCl ₂	(C ₅ Me ₄ Et) ₂ TiMe ₂
(neomenthylCp) ₂ ZrCl ₂	[O(SiMe ₂ Cp) ₂]TiMe ₂
(C ₅ Me ₄ Et) ₂ ZrCl ₂	[O(SiMe ₂ t-BuCp) ₂]TiMe ₂
[O(SIMe ₂ Cp) ₂]ZrCl ₂	Ind ₂ HfMe ₂
[O(SiMe ₂ t-BuCp) ₂]ZrCl ₂	(2-MeInd) ₂ HfMe ₂
Ind ₂ TiCl ₂	(neomenthylCp) ₂ HfMe ₂
(2-MeInd),TICI	(C ₅ Me ₄ Et) ₂ HfMe ₂
(neomenthylCp) ₂ TiCl ₂	[O(SIMe ₂ Cp) ₂]HfMe ₂
(C ₅ Me ₄ Et) ₂ TiCl ₂	[O(SiMe ₂ t-BuCp) ₂]HfMe ₂
[En(Ind) ₂]ZrCl ₂	[Me ₂ Si(IndFlu)]HfCl ₂
[En(Ind) ₂]HfCl ₂	[Me ₂ Si(IndFlu)]TiCl ₂
[En(Ind) ₂]TiCl ₂	[Me ₂ Si(IndFlu)]ZrMe ₂
[En(Ind) ₂]ZrMe ₂	[Me ₂ Si(IndFlu)]HfMe ₂
[En(Ind) ₂]HfMe ₂	[Me ₂ SI(IndFlu)]TiMe ₂
[En(Ind) ₂]TiMe ₂	[Me ₂ Si(IndFlu)]ZrCIMe
[En(Ind) ₂]ZrClMe	[Me ₂ Si(IndFlu)]HfCIMe
[En(Ind) ₂]HfClMe	[Me ₂ Si(IndFlu)]TiClMe
[En(Ind) ₂)TiClMe	[Bz ₂ Si(IndFlu)]ZrCl ₂
[Me ₂ Si(Ind) ₂]ZrCl ₂	[Bz ₂ Si(IndFlu)]HfCl ₂
[Me ₂ Si(Ind) ₂]HfCl ₂	[Bz ₂ Si(IndFlu)]TICl ₂
[Me ₂ Si(Ind) ₂]TiCl ₂	[Bz ₂ Si(IndFlu)]ZrMe ₂
[Me ₂ Si(Ind) ₂]ZrMe ₂	[Bz ₂ Si(IndFlu)]HfMe ₂
[Me ₂ Si(Ind) ₂]HfMe ₂	[Bz ₂ Si(IndFlu)]TiMe ₂
[Me ₂ Si(Ind) ₂]TiMe ₂	[Bz ₂ Si(IndFlu)]ZrClMe
[Me ₂ Si(Ind) ₂]ZrClMe	[Bz ₂ Si(IndFlu)]HfClMe
[Me ₂ Si(Ind) ₂]HfClMe	[Bz ₂ Sl(IndFlu)]TiClMe
[Me ₂ Si(Ind) ₂]TiClMe	[Bimethyl Naphtyl(IndFlu)]ZrCl ₂

Table 1. (continued)

	lable 1. (i	continued)
	List of Metallocenes	
	[Bz ₂ Si(Ind) ₂]ZrCl ₂	[Bimethyl Naphtyl(IndFlu)]HfCl ₂
	[Bz ₂ Si(Ind) ₂]HfCl ₂	[Birnethyl Naphtyl(IndFlu)]TiCl ₂
	[Bz ₂ Si(Ind) ₂]TiCl ₂	[Bimethyl Naphtyl(IndFlu)]ZrMe ₂
	[Bz ₂ Sl(Ind) ₂]ZrMe ₂	[Birnethyl Naphtyl(IndFlu)]HfMe2
	[Bz ₂ Si(Ind) ₂]HfMe ₂	[Birnethyl Naphtyl(IndFlu)]TiMe2
	[Bz ₂ Si(Ind) ₂]TiMe ₂	[Bimethyl Naphtyl(IndFlu)]ZrCIMe
	[Bz ₂ Sl(Ind) ₂]ZrCIMe	[Bimethyl Naphtyl(IndFlu)]HfCIMe
	[Bz ₂ Si(Ind) ₂]HfCIMe	[Bimethyl Naphtyl(IndFlu)]TICIMe
	[Bz ₂ Si(Ind) ₂]TICIMe	[En(BenzindFlu)]ZrCl ₂
	[Bimethyl Naphtyl(Ind) ₂]ZrCl ₂	[En(BenzindFlu)]HfCl ₂
	[Bimethyl Naphtyl(Ind) ₂]HfCl ₂	[En(BenzindFlu)]TiCl ₂
	[Bimethyl Naphtyl(Ind) ₂]TiCl ₂	[En(BenzindFlu)]ZrMe ₂
	[Bimethyl Naphtyl(Ind) ₂]ZrMe ₂	[En(BenzindFlu)]HfMe ₂
	[Bimethyl Naphtyl(Ind) ₂]HfMe ₂	[En(BenzindFlu)]TiMe ₂
	[Bimethyl Naphtyl(Ind) ₂]TiMe ₂	[En(BenzindFlu)]ZrClMe
	[Bimethyl Naphtyl(Ind) ₂]ZrCIMe	[En(BenzindFlu)]HfClMe
	[Bimethyl Naphtyl(Ind) ₂]HfCIMe	[En(BenzindFlu)]TiClMe
	[Bimethyl Naphtyl(Ind) ₂]TiClMe	[Me ₂ Si(BenzIndFlu)]ZrCl ₂
	[En(IndFlu)]ZrCl ₂	[Me ₂ Si(BenzindFlu)HfCl ₂
	[En(IndFlu)]HfCl ₂	[Me ₂ Si(BenzIndFlu)]TiCl ₂
	[En(IndFlu)]TiCl ₂	[Me ₂ SI(BenzindFlu)]ZrMe ₂
	[En(IndFlu)]ZrMe ₂	[Me ₂ Si(BenzindFlu)]HfMe ₂
	[En(IndFlu)]HfMe ₂	[Me ₂ Si(BenzIndFlu)]TiMe ₂
	[En(IndFlu)]TiMe ₂	[Me ₂ Si(BenzIndFlu)]ZrCIMe
	[En(IndFlu)]ZrCIMe	[Me ₂ Si(BenzIndFlu)]HfClMe
	[En(IndFlu)]HfClMe	[Me ₂ Si(BenzIndFlu)]TiCIMe
	[En(IndFlu)]TiClMe	[Bz ₂ Si(BenzindFlu)]ZrCl ₂
	[Me ₂ Si(indFlu)]ZrCl ₂	[Bz ₂ Si(BenzindFlu)]HfCl ₂
	[Bz ₂ Si(BenzIndFlu)]TiCl ₂	[Bimethyl Naphtyl(IndCp)]ZrMe ₂
	[Bz ₂ Si(BenzIndFlu)]ZrMe ₂	[Birnethyl Naphtyl(IndCp)]HfMe ₂
	[Bz ₂ Si(BenzindFlu)]HfMe ₂	[Bimethyl Naphtyl(IndCp)]TiMe ₂
	[Bz ₂ Si(BenzindFlu)]TiMe ₂	[Birnethyl Naphtyl(IndCp)]ZrCIMe
	[Bz ₂ Si(BenzindFlu)]ZrCiMe	
	[Bz ₂ Si(BenzindFlu)]HfClMe	[Birnethyl Naphtyl(IndCp)]HfCIMe [Birnethyl Naphtyl(IndCp)]TiCIMe
	[Bz ₂ Si(BenzindFlu)]TiClMe	[En(FluCp)]ZrCl ₂
	[Bimethyl Naphtyl(BenzindFlu)]ZrCl ₂	[En(FluCp)]HfCl ₂
	[Bimethyl Naphtyl(BenzindFlu)]HfCl ₂	[En(FluCp)]TICl ₂
	[Bimethyl Naphtyl(BenzindFlu)]TiCl ₂	[En(FluCp)]IrCl ₂ [En(FluCp)]ZrMe ₂
	[Birnethyl	
	Naphtyl(BenzindFlu)[ZrMe ₂	[En(FluCp)]HfMe ₂
	[Bimethyl	[En(FluCp)]TiMe ₂
	Naphtyl(BenzindFlu)]HfMe ₂	[En(FluCp)]ZrClMe
	[Blmethyl	[En(FluCp)]HfCIMe
- 1	Naphtyl(BenzindFlu)]TiMe ₂	[En(FluCp)]TICIMe
	[Bimethyl	[Me ₂ Si(FluCp)]ZrCl ₂
		[Me ₂ Si(FluCp)]HfCl ₂
	Naphtyl(BenzindFlu)]ZrCiMe	[Me ₂ Si(FluCp)]TiCl ₂
	[Bimethyl	[Me ₂ Si(FluCp)]ZrMe ₂
	Naphtyl(BenzindFlu)]HfCiMe	[Me ₂ Si(FluCp)]HfMe ₂
1	[Bimethyl	[Me ₂ Si(FluCp)]TiMe ₂
	Naphtyl(BenzindFlu)]TiCiMe	[Me ₂ Si(FluCp)]ZrClMe

Table 1. (continued)

	Table 1. (4	continuea)
	List of Metallocenes	
	[En(IndCp)]ZrCl ₂	[Me ₂ Si(FluCp)]HfClMe
	[En(IndCp)]HfCl ₂	[Me ₂ Si(FluCp)]TiClMe
	[En(IndCp)]TiCl ₂	[Bz ₂ Si(FluCp)]ZrCl ₂
	[En(IndCp)]ZrMe ₂	[Bz ₂ SI(FluCp)]HfCl ₂
	[En(IndCp)]HfMe ₂	[Bz ₂ Si(FluCp)]TiCl ₂
	[En(IndCp)]TiMe ₂	[Bz ₂ Si(FluCp)]ZrMe ₂
	[En(IndCp)]ZrCIMe	[Bz ₂ Si(FluCp)]HfMe ₂
	[En(IndCp)]HfCIMe	[Bz ₂ Si(FluCp)]TiMe ₂
	[En(IndCp)]TiClMe	[Bz ₂ SI(FluCp)]ZrClMe
	[Me ₂ SI(IndCp)]ZrCl ₂	[Bz ₂ Si(FluCp)]HfClMe
	[Me ₂ Si(IndCp)]HfCl ₂	[Bz ₂ Si(FluCp)]TiClMe
	[Me ₂ Si(IndCp)]TiCl ₂	[Bimethyl Naphtyl(FluCp)]ZrCl ₂
	[Me ₂ Si(IndCp)]ZrMe ₂	[Birnethyl Naphtyl(FluCp)]HfCl ₂
	[Me ₂ Si(IndCp)]HfMe ₂	[Bimethyl Naphtyl(FluCp)]TiCl ₂
	[Me ₂ Si(IndCp)]TiMe ₂	[Bimethyl Naphtyl(FluCp)]ZrMe ₂
	[Me ₂ Si(IndCp)]ZrCIMe	[Bimethyl Naphtyl(FluCp)]HfMe2
	[Me ₂ Si(IndCp)]HfCIMe	[Bimethyl Naphtyl(FluCp)]TiMe2
	[Me ₂ Si(IndCp)]TiClMe	[Bimethyl Naphtyl(FluCp)]ZrClMe
	[Bz ₂ Si(IndCp)]ZrCl ₂	[Bimethyl Naphtyl(FluCp)]HfClMe
	[Bz ₂ Si(IndCp)]HfCl ₂	[Bimethyl Naphtyl(FluCp)]TiClMe
	[Bz ₂ Si(IndCp)]TiCl ₂	[En(IndH ₄) ₂]ZrCl ₂
	[Bz ₂ Si(IndCp)]ZrMe ₂	[En(IndH ₄) ₂]HfCl ₂
	[Bz ₂ Si(IndCp)]HfMe ₂	[En(IndH ₄) ₂]TiCl ₂
	[Bz ₂ Si(IndCp)]TiMe ₂	[En(IndH ₄) ₂]ZrMe ₂
	[Bz ₂ Si(IndCp)]ZrCIMe	[Me ₂ Si(Flu) ₂]TiMe ₂
	[Bz ₂ Si(IndCp)]HfCIMe	[Me ₂ Si(Flu) ₂]ZrClMe
	[Bz ₂ Si(IndCp)]TiClMe	[Me ₂ Si(Flu) ₂]HfClMe
	[Bimethyl Naphtyl(IndCp)]ZrCl ₂	[Me ₂ SI(Flu) ₂]TiCiMe
	[Bimethyl Naphtyl(IndCp)]HfCl ₂	[Bz ₂ Si(Flu) ₂]ZrCl ₂
	[Bimethyl Naphtyl(IndCp)]TiCl ₂	[Bz ₂ Si(Flu) ₂]HfCl ₂
	[En(IndH ₄) ₂]HfMe ₂	[Bz ₂ Si(Fiu) ₂]TiCl ₂
	[En(IndH ₄) ₂]TiMe ₂	[Bz ₂ Si(Fiu) ₂]ZrMe ₂
	[En(IndH ₄) ₂]ZrClMe	[Bz ₂ Si(Flu) ₂]HfMe ₂
ı	[En(IndH ₄) ₂]HfClMe	[Bz ₂ Si(Flu) ₂]TiMe ₂
ı	[En(IndH ₄) ₂]TiCIMe	[Bz ₂ Si(Flu) ₂]ZrClMe
	[Bz ₂ Si(IndH ₄) ₂]ZrCl ₂	[Bz ₂ SI(Flu) ₂]HfClMe
į	[Bz ₂ Si(IndH ₄) ₂]HfCl ₂	[Bz ₂ Si(Flu) ₂]TiClMe
	[Bz ₂ Si(IndH ₄) ₂]TiCl ₂	[Bimethyl Naphtyl(Flu) ₂]ZrCl ₂
	[Bz ₂ Si(IndH ₄) ₂]ZrMe ₂	[Bimethyl Naphtyl(Flu) ₂]HfCl ₂
	[Bz ₂ Si(IndH ₄) ₂]HfMe ₂	[Bimethyl Naphtyl(Flu) ₂]TiCl ₂
	[Bz ₂ Si(IndH ₄) ₂]TiMe ₂	[Bimethyl Naphtyl(Flu) ₂]ZrMe ₂
	[Bz ₂ Sl(IndH ₄) ₂]ZrClMe	[Blmethyl Naphtyl(Flu) ₂]HfMe ₂
	[Bz ₂ Si(IndH ₄) ₂]HfClMe	[Bimethyl Naphtyl(Flu)2]TIMe2
	[Bz ₂ Si(IndH ₄) ₂]TiCIMe	[Birnethyl Naphtyl(Flu)2]ZrClMe
1	[Me ₂ Si(IndH ₄) ₂]ZrCl ₂	[Bimethyl Naphtyl(Flu)2]HfClMe
	[Me ₂ Si(IndH ₄) ₂]HfCl ₂	[Bimethyl Naphtyl(Flu)2]TiClMe
	[Me ₂ Si(IndH ₄) ₂]TiCl ₂	[En(2,4,7 Me ₃ Ind) ₂ ZrCl ₂
	[Me ₂ Si(IndH ₄) ₂]ZrMe ₂	[En(IndH ₄) ₂]ZrCl ₂
1	[Me ₂ Si(IndH ₄) ₂]HfMe ₂	[Me ₂ Si (2,4,7 Me ₃ Ind) ₂ ZrCl ₂
	[Me ₂ Si(IndH ₄) ₂]TiMe ₂	[Me ₂ Si (IndH ₄) ₂]ZrCl ₂
		7767 6

Table 1. (continued)

	continued)
List of Metallocenes	
[Me ₂ Si(IndH ₄) ₂]ZrCIMe	[Me ₂ Si(Ind)2]ZrCl ₂
[Me ₂ Si(IndH ₄) ₂]HfClMe	[Ph ₂ Si(Ind) ₂]ZrCl ₂
[Me ₂ Si(IndH ₄) ₂]TiClMe	[Bz ₂ Si(Ind) ₂]ZrCl ₂
[Bimethyl Naphtyl(IndH ₄) ₂]ZrCl ₂	[Me ₂ Si(2,4,7 Me-3-ind) ₂ ZrCi ₂
[Birnethyl Naphtyl(IndH ₄) ₂]HfCl ₂	[Me ₂ Si(IndH ₄) ₂]ZrCl ₂
[Bimethyl Naphtyl(IndH ₄) ₂]TiCl ₂	[Me ₂ Si(2-Me-4,6-I-PrInd) ₂]ZrCl ₂
[Bimethyl Naphtyl(IndH ₄) ₂]ZrMe ₂	[Me ₂ SI(2Me 4Phind) ₂]ZrCl ₂
[Bimethyl Naphtyl(IndH ₄) ₂]HfMe ₂	[Me ₂ SI(2Me4,4Benzoind) ₂]ZrCl ₂
[Bimethyl Naphtyl(IndH ₄) ₂]TiMe ₂	[Me ₂ SI(2,4,7 Me-3-Ind) ₂ ZrCl ₂
[Bimethyl Naphtyl(IndH ₄) ₂]ZrCIMe	[Bz ₂ Si(IndH ₄) ₂]ZrCl ₂
[Bimethyl Naphtyl(IndH ₄) ₂]HfCIMe	[Bz 2Si(2-Me-4,6-I-PrInd)2]ZrCl2
[Bimethyl Naphtyl(IndH ₄) ₂]TiClMe	[Bz 2Si(2Me 4PhInd)2]ZrCl2
[En(Flu) ₂]ZrCl ₂	[Bz 2Si(2Me4,4BenzoInd)2]ZrCl2
[En(Flu) ₂]HfCl ₂	[Ph ₂ C(Ind)(Cp)]ZrCl ₂
[En(Flu)2]TICI2	[Me ₂ C(Ind)(Cp)]ZrCl ₂
[En(Flu) ₂]ZrMe ₂	[Me ₂ C(Ind)(3-MeCp)]ZrCl ₂
[En(Flu) ₂]HfMe ₂	[Ph ₂ C(Flu)(Cp)]ZrCl ₂
[En(Flu) ₂]TiMe ₂	[Me ₂ C(Flu)(Cp)]ZrCl ₂
[En(Flu) ₂]ZrClMe	[Me ₂ C(Flu)(Cp)]HfCl ₂
[En(Flu) ₂]HfClMe	Et(Ind) ₂ ZrCl ₂
[En(Flu) ₂]TiCiMe	Me ₂ Si(Ind) ₂ ZrCl ₂
[Me ₂ SI(Flu) ₂]ZrCl ₂	[Me ₂ Si(2-Me-4,6-I-Prind) ₂]TiCl ₂
[Me ₂ Si(Flu) ₂]HfCl ₂	[Me ₂ Si(2Me 4PhInd) ₂]TiCl ₂
[Me ₂ Si(Flu) ₂]TiCl ₂	[Me ₂ Si(2Me4,4BenzoInd) ₂]TiCl ₂
[Me ₂ Si(Flu) ₂]ZrMe ₂	[Me ₂ SI(2,4,7 Me-3-Ind) ₂ TICl ₂
[Me ₂ Si(Flu) ₂]HfMe ₂	[Bz ₂ Si(IndH ₄) ₂]TiCl ₂
Me ₂ Si(IndH ₄) ₂ ZrCl ₂	[Bz 2SI(2-Me-4,6-i-Prind)2]TiCl2
Me ₂ Si(2-Meind) ₂ ZrCl ₂	[Bz 2SI(2Me 4Phind)2]TICl2
Me ₂ Si(2-Me-4-iPrInd) ₂ ZrCl ₂	[Bz 2Si(2Me4,4Benzolnd)2]TiCl2
Me ₂ Si (2,4-Me ₂ Cp) ₂ ZrCl ₂	[Ph ₂ C(Ind)(Cp)]TiCl ₂
Me ₂ SI(2-Me-4-tBuCp) ₂ ZrCl ₂	
Me ₂ Si(2-Me-4,5 Benzind) ₂ ZrCl ₂	[Me ₂ C(Ind)(Cp)]TICl ₂
	[Me ₂ C(Ind)(3-MeCp)]TICI ₂
Me ₂ Si(2-Me-4-Phind) ₂ ZrCi ₂	[Ph ₂ C(Flu)(Cp)]TICl ₂
Me ₂ Ge(2-Me-4-Phind) ₂ ZrCl ₂	[Me ₂ C(Flu)(Cp)]TICl ₂
Me ₂ Si(2-Me-4-naphthInd) ₂ ZrCl ₂	[Me ₂ C(Flu)(Cp)]HfCl ₂
Bz ₂ Si(Ind) ₂ ZrCl ₂	Et(Ind) ₂ TiCl ₂
Bz ₂ Si(IndH ₄) ₂ ZrCl ₂	Me ₂ Si(Ind) ₂ TiCl ₂
Bz ₂ Si(2-MeInd) ₂ ZrCl ₂	Me ₂ Si(IndH ₄) ₂ TiCl ₂
Bz ₂ Si(2-Me-4-iPrInd) ₂ ZrCl ₂	Me ₂ SI(2-MeInd) ₂ TiCl ₂
Bz ₂ Si (2,4-Me ₂ Cp) ₂ ZrCl ₂	Me ₂ Si(2-Me-4-iPrInd) ₂ TiCl ₂
Bz ₂ Si(2-Me-4-tBuCp) ₂ ZrCl ₂	Me ₂ Si (2,4-Me ₂ Cp) ₂ TiCl ₂
Bz ₂ Si(2-Me-4,5 Benzind) ₂ ZrCl ₂	Me ₂ SI(2-Me-4-tBuCp) ₂ TiCl ₂
Bz ₂ Si(2-Me-4-Phind) ₂ ZrCl ₂	Me ₂ SI(2-Me-4,5 BenzInd) ₂ TiCl ₂
Bz ₂ Ge(2-Me-4-Phind) ₂ ZrCl ₂	Me ₂ Si(2-Me-4-PhInd) ₂ TiCl ₂
Bz ₂ Si(2-Me-4-naphthInd) ₂ ZrCl ₂	Me ₂ Ge(2-Me-4-Phind) ₂ TiCl ₂
Et(IndH ₄) ₂ ZrCl ₂	Me ₂ Si(2-Me-4-naphthind) ₂ TiCl ₂
Et(2-MeInd) ₂ ZrCl ₂	Bz ₂ Si(Ind) ₂ TiCl ₂
Et(2-Me-4-iPrInd) ₂ ZrCl ₂	Bz ₂ Si(IndH ₄) ₂ TiCl ₂
Et(2,4-Me ₂ Cp) ₂ ZrCl ₂	Bz ₂ Si(2-MeInd) ₂ TiCl ₂
Et(2-Me-4-tBuCp) ₂ ZrCl ₂	Bz ₂ Si(2-Me-4-iPrInd) ₂ TiCl ₂

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Table 1. (continued)

List of Metallocenes	
Et(2-Me-4,5 Benzind) ₂ ZrCl ₂	Bz ₂ Si (2,4-Me ₂ Cp) ₂ TiCl ₂
Et(2-Me-4-Phind) ₂ ZrCl ₂	Bz ₂ Si(2-Me-4-tBuCp) ₂ TiCl ₂
Et(2-Me-4-Phind) ₂ ZrCl ₂	Bz ₂ Si(2-Me-4,5 BenzInd) ₂ TiCl ₂
Et(2-Me-4-naphthind) ₂ ZrCl ₂	Bz ₂ Si(2-Me-4-Phind) ₂ TiCl ₂
Et(Ind) ₂ ZrCl ₂	Bz ₂ Ge(2-Me-4-Phind) ₂ TiCl ₂
Et(IndH ₄) ₂ ZrCl ₂	Bz ₂ Si(2-Me-4-naphthInd) ₂ TiCl ₂
Et (2-MeInd) ₂ ZrCl ₂	Et(IndH ₄) ₂ TICl ₂
Et(2-Me-4-iPrind) ₂ ZrCl ₂	Et(2-Meind) ₂ TiCl ₂
Et(2,4-Me ₂ Cp) ₂ ZrCl ₂	Et(2-Me-4-IPrind) ₂ TICl ₂
Et(2-Me-4-tBuCp) ₂ ZrCl ₂	Et(2,4-Me ₂ Cp) ₂ TICl ₂
Et(2-Me-4,5 Benzind) ₂ ZrCl ₂	Et(2-Me-4-tBuCp) ₂ TiCl ₂
Et(2-Me-4-Phind) ₂ ZrCl ₂	Et(2-Me-4,5 Benzind) ₂ TiCl ₂
Et(2-Me-4-Phind) ₂ ZrCl ₂	Et(2-Me-4-Phind) ₂ TiCl ₂
Et(2-Me-4-naphthind) ₂ ZrCl ₂	Et(2-Me-4-Phind) ₂ TiCl ₂
[En(2,4,7 Me ₃ Ind) ₂ TiCl ₂	Et(2-Me-4-naphthInd) ₂ TiCl ₂
[En(IndH ₄) ₂]TiCl ₂	Et(Ind) ₂ TiCl ₂
[Me ₂ Si (2,4,7 Me ₃ Ind) ₂ TiCl ₂	Et(IndH ₄) ₂ TiCl ₂
[Me ₂ SI (IndH ₄) ₂]TICl ₂	Bz ₂ Si(2-Me-4-tBuCp) ₂ HfCl ₂
[Me ₂ SI(Ind)2]TiCl ₂	Bz2SI(2-Me-4,5 Benzind) ₂ HfCl ₂
[Ph ₂ Si(Ind) ₂]TiCl ₂	Bz ₂ Si(2-Me-4-PhInd) ₂ HfCl ₂
[Bz ₂ Si(ind) ₂]TiCl ₂	Bz ₂ Ge(2-Me-4-Phind) ₂ HfCl ₂
[Me ₂ Si(2,4,7 Me-3-Ind) ₂ TiCl ₂	Bz ₂ SI(2-Me-4-naphthInd) ₂ HfCl ₂
[Me ₂ Si(IndH ₄) ₂]TiCl ₂	Et(IndH ₄) ₂ HfCl ₂
Et (2-Melnd),TiCl,	Et(2-Melnd) ₂ HfCl ₂
Et(2-Me-4-iPrInd) ₂ TiCl ₂	Et(2-Me-4-IPrind) ₂ HfCl ₂
Et(2,4-Me ₂ Cp) ₂ TiCl ₂	Et(2,4-Me ₂ Cp) ₂ HfCl ₂
Et(2-Me-4-tBuCp) ₂ TiCl ₂	Et(2-Me-4-tBuCp) ₂ HfCl ₂
Et(2-Me-4,5 Benzind) ₂ TiCl ₂	Et(2-Me-4,5 Benzind) ₂ HfCl ₂
Et(2-Me-4-Phind) ₂ TiCl ₂	Et(2-Me-4-Phind) ₂ HfCl ₂
Et(2-Me-4-Phind)2TiCl2	Et(2-Me-4-Phind) ₂ HfCl ₂
Et(2-Me-4-naphthind)2TiCl2	Et(2-Me-4-naphthind) ₂ HfCl ₂
[En(2,4,7 Me ₃ Ind) ₂ HfCl ₂	Et(Ind) ₂ HfCl ₂
[En(IndH ₄) ₂]HfCl ₂	Et(IndH ₄) ₂ HfCl ₂
[Me ₂ Si (2,4,7 Me ₃ Ind) ₂ HfCl ₂	Et (2-MeInd) ₂ HfCl ₂
[Me ₂ Si (IndH ₄) ₂]HfCl ₂	Et(2-Me-4-iPrind) ₂ HfCl ₂
[Me ₂ SI(Ind)2]HfCl ₂	Et(2,4-Me ₂ Cp) ₂ HfCl ₂
[Ph ₂ Si(Ind) ₂]HfCl ₂	Et(2-Me-4-tBuCp) ₂ HfCl ₂
[Bz ₂ Si(Ind) ₂]HfCl ₂	Et(2-Me-4,5 Benzind) ₂ HfCl ₂
[Me ₂ Si(2,4,7 Me-3-Ind) ₂ HfCl ₂	Et(2-Me-4-Phind) ₂ HfCl ₂
[Me ₂ Si(IndH ₄) ₂]HfCl ₂	Et(2-Me-4-Phind) ₂ HfCl ₂
[Me ₂ Si(2-Me-4,6-I-PrInd) ₂]HfCl ₂	Et(2-Me-4-naphthind) ₂ HfCl ₂
[Me ₂ SI(2Me-4,64FHIIId) ₂]HfCl ₂	
[Me ₂ SI(2Me4,4BenzoInd) ₂]HfCl ₂	[En(2,4,7 Me ₃ ind) ₂ ZrMe ₂ [En(IndH ₄) ₂]ZrMe ₂
[Me ₂ SI(2,4,7 Me-3-Ind) ₂ HfCl ₂	[Me ₂ Si (2,4,7 Me ₃ Ind) ₂ ZrMe ₂
[Bz ₂ Si(IndH ₄) ₂]HfCl ₂	
[Bz ₂ Si(2-Me-4,6-i-Prind) ₂]HfCl ₂	[Me ₂ Si (IndH ₄) ₂]ZrMe ₂
	[Me ₂ Si(Ind)2]ZrMe ₂
[Bz ₂ Si(2Me 4Phind) ₂]HfCl ₂	[Ph ₂ Si(Ind) ₂]ZrMe ₂
[Bz 2Si(2Me4,4BenzoInd)2]HfCl2	[Bz ₂ Si(Ind) ₂]ZrMe ₂
[Ph ₂ C(Ind)(Cp)]HfCl ₂	[Me ₂ Si(2,4,7 Me-3-Ind) ₂ ZrMe ₂
[Me ₂ C(Ind)(Cp)]HfCl ₂	[Me ₂ Si(IndH ₄) ₂]ZrMe ₂

Table 1. (continued)

rabie i.	(continued)
List of Metallocenes	
[Me ₂ C(Ind)(3-MeCp))HfCl ₂	[Me ₂ Si(2-Me-4,6-i-PrInd) ₂]ZrMe ₂
[Ph ₂ C(Flu)(Cp)]HfCl ₂	[Me ₂ Si(2Me 4PhInd) ₂]ZrMe ₂
[Me ₂ C(Flu)(Cp)]HfCl ₂	[Me ₂ Si(2Me4,4BenzoInd) ₂]ZrMe ₂
[Me ₂ C(Flu)(Cp)]HfCl ₂	[Me ₂ Si(2,4,7 Me-3-Ind) ₂ ZrMe ₂
Et(Ind) ₂ HfCl ₂	[Bz ₂ Si(IndH ₄) ₂]ZrMe ₂
Me ₂ Si(Ind) ₂ HfCl ₂	[Bz 2Si(2-Me-4,6-I-Prind)2]ZrMe2
Me ₂ Si(IndH ₄) ₂ HfCl ₂	[Bz 2SI(2Me 4Phind)2]ZrMe2
Me ₂ Si(2-MeInd) ₂ HfCl ₂	[Bz 2Si(2Me4,4Benzolnd)2]ZrMe2
Me ₂ Si(2-Me-4-iPrInd) ₂ HfCl ₂	[Ph ₂ C(Ind)(Cp)]ZrMe ₂
Me ₂ SI (2,4-Me ₂ Cp) ₂ HfCl ₂	[Me ₂ C(Ind)(Cp)]ZrMe ₂
Me ₂ Si(2-Me-4-tBuCp) ₂ HfCl ₂	[Me ₂ C(Ind)(3-MeCp)]ZrMe ₂
Me ₂ Si(2-Me-4,5 Benzind) ₂ HfCl ₂	[Ph ₂ C(Flu)(Cp)]ZrMe ₂
Me ₂ Si(2-Me-4-Phind) ₂ HfCl ₂	[Me ₂ C(Flu)(Cp)]ZrMe ₂
Me ₂ Ge(2-Me-4-Phind) ₂ HfCl ₂	[Me ₂ C(Flu)(Cp)]HfMe ₂
Me ₂ Si(2-Me-4-naphthInd) ₂ HfCl ₂	[Me ₂ Si(2,4,7 Me-3-Ind) ₂ TiMe ₂
Bz ₂ Si(Ind) ₂ HfCl ₂	[Me ₂ Si(IndH ₄) ₂]TiMe ₂ [Me ₂ Si(2-Me-4,6-i-Prind) ₂]TiMe ₂
Bz ₂ Si(IndH ₄) ₂ HfCl ₂	
Bz ₂ Si(2-MeInd) ₂ HfCl ₂	[Me ₂ SI(2Me 4Phind) ₂]TiMe ₂
Bz ₂ Si(2-Me-4-IPrind) ₂ HfCl ₂	[Me ₂ Si(2Me4,4BenzoInd) ₂]TiMe ₂
Bz ₂ Si (2,4-Me ₂ Cp) ₂ HfCl ₂	[Me ₂ Si(2,4,7 Me-3-Ind) ₂ TiMe ₂
Et(Ind) ₂ ZrMe ₂	[Bz ₂ Si(IndH ₄) ₂]TiMe ₂
Me ₂ SI(Ind) ₂ ZrMe ₂	[Bz ₂ Si(2-Me-4,6-I-Prind) ₂]TiMe ₂
Me ₂ Si(IndH ₄) ₂ ZrMe ₂	[Bz ₂ Si(2Me 4PhInd) ₂]TiMe ₂
Me ₂ Si(2-MeInd) ₂ ZrMe ₂	[Bz ₂ Si(2Me4,4Benzolnd) ₂]TiMe ₂
Me ₂ Si(2-Me-4-iPrind) ₂ ZrMe ₂	[Ph ₂ C(Ind)(Cp)]TiMe ₂
Me ₂ Si (2,4-Me ₂ Cp) ₂ ZrMe ₂	[Me ₂ C(Ind)(Cp)]TiMe ₂
Me ₂ Si(2-Me-4-tBuCp) ₂ ZrMe ₂	[Me ₂ C(Ind)(3-MeCp)]TiMe ₂
Me ₂ Si(2-Me-4,5 BenzInd) ₂ ZrMe ₂	[Ph ₂ C(Flu)(Cp)]TiMe ₂
Me ₂ Si(2-Me-4-PhInd) ₂ ZrMe ₂	[Me ₂ C(Flu)(Cp)]TiMe ₂
Me ₂ Ge(2-Me-4-Phind) ₂ ZrMe ₂	[Me ₂ C(Flu)(Cp)]HfMe ₂
Me ₂ Si(2-Me-4-naphthInd) ₂ ZrMe ₂	Et(Ind) ₂ TiMe ₂
Bz ₂ Si(Ind) ₂ ZrMe ₂	Me ₂ Si(Ind) ₂ TiMe ₂
Bz ₂ Si(IndH ₄) ₂ ZrMe ₂	Me ₂ SI(IndH ₄) ₂ TiMe ₂
Bz ₂ Si(2-MeInd) ₂ ZrMe ₂	Me ₂ Si(2-MeInd) ₂ TiMe ₂
Bz ₂ Si(2-Me-4-iPrInd) ₂ ZrMe ₂	Me ₂ Si(2-Me-4-iPrInd) ₂ TiMe ₂
Bz ₂ Si (2,4-Me ₂ Cp) ₂ ZrMe ₂	Me ₂ SI (2,4-Me ₂ Cp) ₂ TiMe ₂
Bz ₂ Si(2-Me-4-tBuCp) ₂ ZrMe ₂	Me ₂ Si(2-Me-4-tBuCp) ₂ TiMe ₂
Bz ₂ Si(2-Me-4,5 BenzInd) ₂ ZrMe ₂	Me ₂ Si(2-Me-4,5 BenzInd) ₂ TIMe ₂
Bz ₂ Si(2-Me-4-PhInd) ₂ ZrMe ₂	Me ₂ Si(2-Me-4-PhInd) ₂ TiMe ₂
Bz ₂ Ge(2-Me-4-Phind) ₂ ZrMe ₂	Me ₂ Ge(2-Me-4-Phind) ₂ TiMe ₂
Bz ₂ Si(2-Me-4-naphthInd) ₂ ZrMe ₂	Me ₂ Si(2-Me-4-naphthind) ₂ TiMe ₂
Et(IndH ₄) ₂ ZrMe ₂	Bz ₂ Si(Ind) ₂ TiMe ₂
Et(2-MeInd) ₂ ZrMe ₂	Bz ₂ SI(IndH ₄) ₂ TiMe ₂
Et(2-Me-4-iPrind) ₂ ZrMe ₂	Bz ₂ Si(2-MeInd) ₂ TiMe ₂
Et(2,4-Me, Cp) ,ZrMe,	Bz ₂ Si(2-Me-4-iPrind) ₂ TiMe ₂
Et(2-Me-4-tBuCp) ₂ ZrMe ₂	Bz ₂ SI (2,4-Me ₂ Cp) ₂ TiMe ₂
Et(2-Me-4,5 Benzind) ₂ ZrMe ₂	Bz ₂ Si(2-Me-4-tBuCp) ₂ TiMe ₂
Et(2-Me-4-Phind) ₂ ZrMe ₂	Bz ₂ Si(2-Me-4,5 BenzInd) ₂ TiMe ₂
Et(2-Me-4-Phind) ₂ ZrMe ₂	Bz ₂ Si(2-Me-4-Phind) ₂ TiMe ₂
Et(2-Me-4-naphthInd) ₂ ZrMe ₂	Bz ₂ Ge(2-Me-4-Phind) ₂ TiMe ₂
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Table 1. (continued)

List of Metallocenes Et(Ind) ₂ ZrMe ₂	Bz ₂ Si(2-Me-4-naphthInd) ₂ TiMe ₂
Et(IndH ₄) ₂ ZrMe ₂	Et(IndH ₄) ₂ TiMe ₂
Et (2-Meind) ₂ ZrMe ₂	Et(2-MeInd) ₂ TiMe ₂
Et(2-Me-4-IPrind) ₂ ZrMe ₂	Et(2-Me-4-iPrind) ₂ TiMe ₂
Et(2,4-Me ₂ Cp) ₂ ZrMe ₂	Et(2,4-Me ₂ Cp) ₂ TiMe ₂
Et(2-Me-4-tBuCp) ₂ ZrMe ₂	Et(2-Me-4-tBuCp) ₂ TiMe ₂
Et(2-Me-4,5 Benzind) ₂ ZrMe ₂	Et(2-Me-4,5 Benzind) ₂ TiMe ₂
Et(2-Me-4-Phind) ₂ ZrMe ₂	Et(2-Me-4-Phind) ₂ TiMe ₂
Et(2-Me-4-Phind) ₂ ZrMe ₂	Et(2-Me-4-Phind) ₂ TiMe ₂
Et(2-Me-4-naphthind) ₂ ZrMe ₂	Et(2-Me-4-naphthind) ₂ TiMe ₂
[En(2,4,7 Me ₃ Ind) ₂ TiMe ₂	Bz ₂ Si(2-Me-4-iPrInd) ₂ HfMe ₂
[En(IndH ₄) ₂]TiMe ₂	Bz ₂ Si (2,4-Me ₂ Cp) ₂ HfMe ₂
[Me ₂ Si (2,4,7 Me ₃ Ind) ₂ TiMe ₂	Bz ₂ Si(2-Me-4-tBuCp) ₂ HfMe ₂
[Me ₂ Si (IndH ₄) ₂]TiMe ₂	Bz ₂ Si(2-Me-4,5 BenzInd) ₂ HfMe ₂
[Me ₂ Si(Ind) ₂]TiMe ₂	Bz ₂ Si(2-Me-4-PhInd) ₂ HfMe ₂
[Ph ₂ Si(Ind) ₂]TiMe ₂	Bz ₂ Ge(2-Me-4-Phind) ₂ HfMe ₂
[Bz ₂ Si(Ind) ₂]TiMe ₂	Bz ₂ Si(2-Me-4-naphthInd) ₂ HfMe ₂
Et(Ind) ₂ TiMe ₂	Et(IndH ₄) ₂ HfMe ₂
Et(IndH ₄) ₂ TiMe ₂	Et(2-MeInd) ₂ HfMe ₂
Et (2-MeInd) ₂ TiMe ₂	Et(2-Me-4-iPrind) ₂ HfMe ₂
Et(2-Me-4-iPrInd) ₂ TiMe ₂	Et(2,4-Me ₂ Cp) ₂ HfMe ₂
Et(2,4-Me ₂ Cp) ₂ TiMe ₂	Et(2-Me-4-tBuCp) ₂ HfMe ₂
Et(2-Me-4-tBuCp) ₂ TiMe ₂	Et(2-Me-4,5 Benzind) ₂ HfMe ₂
Et(2-Me-4,5 Benzind) ₂ TiMe ₂	Et(2-Me-4-Phind) ₂ HfMe ₂
Et(2-Me-4-Phind) ₂ TiMe ₂	Et(2-Me-4-Phind) ₂ HfMe ₂
Et(2-Me-4-Phind) ₂ TiMe ₂	Et(2-Me-4-naphthind) ₂ HfMe ₂
Et(2-Me-4-naphthind) ₂ TiMe ₂	Et(Ind) ₂ HfMe ₂
[En(2,4,7 Me ₃ Ind) ₂ HfMe ₂	Et(IndH ₄) ₂ HfMe ₂
[En(IndH ₄) ₂]HfMe ₂	Et(2-Meind) ₂ HfMe ₂
[Me ₂ Si(2,4,7 Me ₃ Ind) ₂ HfMe ₂	Et(2-Me-4-IPrind) ₂ HfMe ₂
[Me ₂ SI (IndH ₄) ₂]HfMe ₂	Et(2,4-Me ₂ Cp) ₂ HfMe ₂
[Me ₂ SI(Ind)2]HfMe ₂	Et(2-Me-4-tBuCp) ₂ HfMe ₂
[Ph ₂ Si(Ind) ₂]HfMe ₂	Et(2-Me-4,5 Benzind) ₂ HfMe ₂
[Bz ₂ Si(Ind) ₂]HfMe ₂	Et(2-Me-4-Phind) ₂ HfMe ₂
[Me ₂ Si(2,4,7 Me-3-Ind) ₂ HfMe ₂	Et(2-Me-4-Phind) ₂ HfMe ₂
[Me ₂ SI(IndH ₄) ₂]HfMe ₂	Et(2-Me-4-naphthind) ₂ HfMe ₂
[Me ₂ Si(2-Me-4,6-i-PrInd) ₂]HfMe ₂	[En(2,4,7 Me ₃ Ind) ₂ ZrCiMe
[Me ₂ Si(2Me 4Phind) ₂]HfMe ₂	[En(IndH ₄) ₂]ZrClMe
Me ₂ Si(2Me4,4Benzolnd) ₂]HfMe ₂	[Me ₂ Si(2,4,7 Me ₃ Ind) ₂ ZrClMe
Me ₂ Si(2,4,7 Me-3-Ind) ₂ HfMe ₂	[Me ₂ Si (IndH ₄) ₂]ZrClMe
[Bz ₂ Si(IndH ₄) ₂]HfMe ₂	[Me ₂ Si(Ind) ₂]ZrClMe
Bz 2Si(2-Me-4,6-i-PrInd)2]HfMe2	[Ph ₂ SI(Ind) ₂]ZrCIMe
[Bz ₂ SI(2Me 4Phind) ₂]HfMe ₂	[Bz ₂ Si(ind) ₂]ZrCiMe
Bz 2Si(2Me4,4BenzoInd)2]HfMe2	[Me ₂ Si(2,4,7 Me-3-Ind) ₂ ZrCIMe
Ph ₂ C(Ind)(Cp)]HfMe ₂	[Me ₂ SI(IndH ₄) ₂]ZrCIMe
Me ₂ C(Ind)(Cp)]HfMe ₂	[Me ₂ Si(2-Me-4,6-i-Prind) ₂]ZrClMe
Me ₂ C(Ind)(3-MeCp)]HfMe ₂	[Me ₂ Si(2Me 4PhInd) ₂]ZrClMe
Ph ₂ C(Flu)(Cp)]HfMe ₂	[Me ₂ Si(2Me4,4Benzolnd) ₂]ZrClMe
[Me ₂ C(Flu)(Cp)]HfMe ₂	[Me ₂ Si(2,4,7 Me-3-Ind) ₂ ZrCiMe
[Me ₂ C(Flu)(Cp)]HfMe ₂	[Bz ₂ Si(IndH ₄) ₂]ZrCIMe

Table 1. (continued)

List of Metallocenes	
Et(Ind) ₂ HfMe ₂	[Bz 2Si(2-Me-4,6-i-Prind)2]ZrCiMe
Me ₂ Si(Ind) ₂ HfMe ₂	[Bz ₂ Si(2Me 4PhInd) ₂]ZrCIMe
Me ₂ Si(IndH ₄) ₂ HfMe ₂	[Bz 2Si(2Me4,4BenzoInd)2]ZrCIMe
Me ₂ Si(2-MeInd) ₂ HfMe ₂	[Ph ₂ C(Ind)(Cp)]ZrCIMe
Me ₂ Si(2-Me-4-iPrInd) ₂ HfMe ₂	[Me ₂ C(Ind)(Cp)]ZrCIMe
Me ₂ Si (2,4-Me ₂ Cp) ₂ HfMe ₂	[Me ₂ C(Ind)(3-MeCp)]ZrCIMe
Me ₂ SI(2-Me-4-tBuCp) ₂ HfMe ₂	[Ph ₂ C(Flu)(Cp)]ZrClMe
Me2SI(2-Me-4,5 Benzind)2HfMe2	[Me ₂ Si(Ind)2]TiCIMe
Me ₂ Si(2-Me-4-Phind) ₂ HfMe ₂	[Ph ₂ Si(ind) ₂]TiCiMe
Me ₂ Ge(2-Me-4-Phind) ₂ HfMe ₂	[Bz ₂ Si(Ind) ₂]TiClMe
Me ₂ Si(2-Me-4-naphthInd) ₂ HfMe ₂	[Me ₂ Si(2,4,7 Me-3-Ind) ₂ TiClMe
Bz ₂ Si(Ind) ₂ HfMe ₂	[Me ₂ SI(IndH ₄) ₂]TiCIMe
Bz ₂ Si(IndH ₄) ₂ HfMe ₂	[Me ₂ Si(2-Me-4,6-I-Prind) ₂]TICIMe
Bz ₂ Si(2-MeInd) ₂ HfMe ₂	[Me ₂ Si(2Me 4PhInd) ₂)TiClMe
[Me ₂ C(Flu)(Cp)]ZrCIMe	[Me ₂ Si(2Me4,4Benzolnd) ₂]TiClMe
[Me ₂ C(Flu)(Cp)]HfClMe	[Me ₂ Si(2,4,7 Me-3-Ind) ₂ TiClMe
Et(Ind) ₂ ZrCIMe	[Bz ₂ Si(IndH ₄) ₂]TiCIMe
Me ₂ SI(Ind) ₂ ZrCIMe	[Bz 2Sl(2-Me-4,6-I-Prind)2]TiCIMe
Me ₂ SI(IndH ₄) ₂ ZrCIMe	[Bz 2SI(2Me 4Phind)2]TICIMe
Me ₂ Si(2-MeInd) ₂ ZrCIMe	[Bz 2SI(2Me4,4BenzoInd)2]TiCIMe
Me ₂ SI(2-Me-4-IPrind) ₂ ZrCIMe	[Ph ₂ C(Ind)(Cp)]TICIMe
Me ₂ SI (2,4-Me ₂ Cp) ₂ ZrCIMe	[Me ₂ C(Ind)(Cp)]TICIMe
Me ₂ Si(2-Me-4-tBuCp) ₂ ZrClMe	
	[Me ₂ C(Ind)(3-MeCp)]TiCIMe
Me ₂ Si(2-Me-4,5 BenzInd) ₂ ZrCIMe Me ₂ Si(2-Me-4-PhInd) ₂ ZrCIMe	[Ph ₂ C(Flu)(Cp)]TiClMe
Me ₂ Ge(2-Me-4-Phind) ₂ ZrCiMe	[Me ₂ C(Flu)(Cp)]TiCiMe
	[Me ₂ C(Flu)(Cp)]HfClMe
Me ₂ SI(2-Me-4-naphthInd) ₂ ZrCIMe Bz ₂ SI(Ind) ₂ ZrCIMe	Et(Ind) ₂ TICIMe
	Me ₂ Si(Ind) ₂ TiCIMe
Bz ₂ Si(IndH ₄) ₂ ZrCIMe	Me ₂ Si(IndH ₄) ₂ TiClMe
Bz ₂ Si(2-MeInd) ₂ ZrClMe	Me ₂ Si(2-MeInd) ₂ TiClMe
Bz ₂ Si(2-Me-4-IPrind) ₂ ZrCIMe	Me ₂ Si(2-Me-4-IPrind) ₂ TiCiMe
Bz ₂ Si (2,4-Me ₂ Cp) ₂ ZrClMe	Me ₂ Si (2,4-Me ₂ Cp) ₂ TICIMe
Bz ₂ Si(2-Me-4-tBuCp) ₂ ZrClMe	Me ₂ Si(2-Me-4-tBuCp) ₂ TiClMe
Bz ₂ Si(2-Me-4,5 BenzInd) ₂ ZrCIMe	Me ₂ Si(2-Me-4,5 BenzInd) ₂ TiClMe
Bz ₂ Si(2-Me-4-PhInd) ₂ ZrClMe	Me ₂ Si(2-Me-4-PhInd) ₂ TiCIMe
Bz ₂ Ge(2-Me-4-PhInd) ₂ ZrCIMe	Me ₂ Ge(2-Me-4-Phind) ₂ TICIMe
Bz ₂ Si(2-Me-4-naphthInd) ₂ ZrCIMe	Me ₂ Si(2-Me-4-naphthind) ₂ TICIMe
Et(IndH ₄) ₂ ZrCIMe	Bz ₂ Si(Ind) ₂ TiCIMe
Et(2-MeInd) ₂ ZrCIMe	Bz ₂ Si(IndH ₄) ₂ TiClMe
Et(2-Me-4-iPrInd) ₂ ZrCIMe	Bz ₂ Si(2-MeInd) ₂ TiCIMe
Et(2,4-Me ₂ Cp) ₂ ZrClMe	Bz ₂ Si(2-Me-4-iPrInd) ₂ TiCIMe
Et(2-Me-4-tBuCp) ₂ ZrCIMe	Bz ₂ Si (2,4-Me ₂ Cp) ₂ TiClMe
Et(2-Me-4,5 Benzind) ₂ ZrCiMe	Bz ₂ Si(2-Me-4-tBuCp) ₂ TICIMe
Et(2-Me-4-Phind) ₂ ZrCiMe	Bz ₂ Si(2-Me-4,5 BenzInd) ₂ TICIMe
Et(2-Me-4-Phind) ₂ ZrCiMe	Bz ₂ Si(2-Me-4-Phind) ₂ TiClMe
Et(2-Me-4-naphthind) ₂ ZrClMe	Bz ₂ Ge(2-Me-4-Phind) ₂ TiClMe
Et(Ind) ₂ ZrCIMe	Bz ₂ Si(2-Me-4-naphthInd) ₂ TiClMe
Et(IndH ₄) ₂ ZrCIMe	Et(IndH ₄) ₂ TICIMe
Et (2-MeInd) ₂ ZrClMe	Et(2-MeInd) ₂ TiCIMe
Et(2-Me-4-iPrInd) ₂ ZrCIMe	Et(2-Me-4-iPrInd) ₂ TiCIMe

Table 1. (continued)

List of Metallocenes	
Et(2,4-Me ₂ Cp) ₂ ZrClMe	Et(2,4-Me ₂ Cp) ₂ TiClMe
Et(2-Me-4-tBuCp) ₂ ZrClMe	Et(2-Me-4-tBuCp) ₂ TiCIMe
Et(2-Me-4,5 Benzind) ₂ ZrCIMe	Et(2-Me-4,5 Benzind) ₂ TiClMe
Et(2-Me-4-Phind) ₂ ZrClMe	Me ₂ Si(2-Me-4-naphthind) ₂ HfClMe
Et(2-Me-4-Phind) ₂ ZrClMe	Bz ₂ Si(Ind) ₂ HfCIMe
Et(2-Me-4-naphthind) ₂ ZrCIMe	Bz ₂ Si(indH ₄) ₂ HfClMe
[En(2,4,7 Me ₃ Ind) ₂ TiClMe	Bz ₂ Sl(2-MeInd) ₂ HfCIMe
[En(IndH ₄) ₂]TiClMe	Bz ₂ Si(2-Me-4-iPrind) ₂ HfClMe
[Me ₂ Si (2,4,7 Me ₃ Ind) ₂ TiClMe	
[Me ₂ SI (IndH ₄) ₂]TICIMe	Bz ₂ Si (2,4-Me ₂ Cp) ₂ HfClMe Bz ₂ Sl(2-Me-4-tBuCp) ₂ HfClMe
Et(2-Me-4-Phind) ₂ TiClMe	
	Bz ₂ Si(2-Me-4,5 BenzInd) ₂ HfClMe
Et(2-Me-4-Phind) ₂ TiClMe	Bz ₂ Si(2-Me-4-PhInd) ₂ HfCIMe
Et(2-Me-4-naphthind) ₂ TiClMe	Bz ₂ Ge(2-Me-4-Phind) ₂ HfClMe
Et(Ind) ₂ TiCIMe	Bz ₂ Si(2-Me-4-naphthInd) ₂ HfCIMe
Et(IndH ₄) ₂ TICIMe	Et(IndH ₄) ₂ HfClMe
Et (2-MeInd) ₂ TICIMe	Et(2-MeInd) ₂ HfCIMe
Et(2-Me-4-iPrInd) ₂ TiClMe	Et(2-Me-4-iPrInd) ₂ HfCIMe
Et(2,4-Me ₂ Cp) ₂ TiClMe	Et(2,4-Me ₂ Cp) ₂ HfClMe
Et(2-Me-4-tBuCp) ₂ TiClMe	Et(2-Me-4-tBuCp) ₂ HfClMe
Et(2-Me-4,5 Benzind) ₂ TiClMe	Et(2-Me-4,5 Benzind) ₂ HfCIMe
Et(2-Me-4-Phind) ₂ TiClMe	Et(2-Me-4-Phind) ₂ HfCiMe
Et(2-Me-4-Phind) ₂ TiClMe	Et(2-Me-4-Phind) ₂ HfClMe
Et(2-Me-4-naphthInd) ₂ TiCIMe	Et(2-Me-4-naphthind) ₂ HfCIMe
[En(2,4,7 Me ₃ Ind) ₂ HfCIMe	Et(Ind) ₂ HfCIMe
[En(IndH ₄) ₂]HfCIMe	Et(IndH ₄) ₂ HfCIMe
[Me ₂ Si (2,4,7 Me ₃ Ind) ₂ HfCIMe	Et (2-Melnd) ₂ HfClMe
[Me ₂ Si (IndH ₄) ₂]HfClMe	Et(2-Me-4-iPrInd) ₂ HfCIMe
[Me ₂ Si(Ind) ₂]HfCIMe	Et(2,4-Me ₂ Cp) ₂ HfClMe
[Ph ₂ Si(Ind) ₂]HfCIMe	Et(2-Me-4-tBuCp) ₂ HfCiMe
[Bz ₂ Si(Ind) ₂]HfCIMe	Et(2-Me-4,5 Benzind) ₂ HfCiMe
[Me ₂ SI(2,4,7 Me-3-Ind) ₂ HfCIMe	Et(2-Me-4-Phind) ₂ HfClMe
[Me ₂ Si(IndH ₄) ₂]HfClMe	Et(2-Me-4-Phind) ₂ HfClMe
[Me ₂ SI(2-Me-4,6-I-Prind) ₂]HfCIMe	Et(2-Me-4-naphthind) ₂ HfCiMe
[Me ₂ Si(2Me 4Phind) ₂]HfClMe	Et(Ind) ₂ HfCIMe
[Me ₂ Si(2Me4,4Benzolnd) ₂]HfClMe	Me ₂ Si(Ind) ₂ HfCIMe
[Me ₂ Si(2,4,7 Me-3-Ind) ₂ HfCIMe	Me ₂ Si(IndH ₄) ₂ HfClMe
[Bz ₂ Si(IndH ₄) ₂]HfCIMe	Me ₂ Si(2-MeInd) ₂ HfCIMe
[Bz ₂ Si(2-Me-4,6-i-Prind) ₂]HfClMe	Me ₂ Si(2-Me-4-iPrInd) ₂ HfCIMe
[Bz 2Si(2Me 4PhInd)2]HfCIMe	Me ₂ Si (2,4-Me ₂ Cp) ₂ HfClMe
[Bz 2Si(2Me4,4Benzolnd)2]HfClMe	Me ₂ Si(2-Me-4-tBuCp) ₂ HfClMe
[Ph ₂ C(Ind)(Cp)]HfCIMe	Me ₂ Ge(2-Me-4-Phind) ₂ HfCiMe
[Me ₂ C(Ind)(Cp)]HfCIMe	Me ₂ Si(2-Me-4,5 BenzInd) ₂ HfClMe
[Me ₂ C(Ind)(3-MeCp)]HfCIMe	Me ₂ Si(2-Me-4-Phind) ₂ HfCiMe
[Ph ₂ C(Flu)(Cp)]HfCIMe	
[Me ₂ C(Flu)(Cp)]HfClMe	
[Me ₂ C(Flu)(Cp)]HfClMe	

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[0084] Preferred metallocene catalysts for ethylene polymerization are (CpR)_ZrX2 catalysts, where Cp is cyclopentadionyl, indemyl, fluorenyl, R is H, Me (methyl), Et (ethyl), Pr (propyl), I-Pr, Bu (butyl), I-Bu, and X is Ct. The metallocene catalyst can be used as part of a catalyst system containing also a co-catalyst which activate the

metallocene in the terpolymerization. Examples of such co-catalysts are alumoxanes such as methyl alumoxane (MAO), ethyl alumoxane (EAO), and isobutylalumoxane.

[0085] Different known methods of adding the cocatalyst can be used, such as:

mixing the metallocene catalyst with the cocatalyst under inert conditions in an inert solvent and bringing the activated complex catalyst formed into the reaction zone prior or continuously during the terpolymerization; or mixing the cocatalyst with solvent provided for the polymerization and introducing further the catalyst to form the catalyst complex prior to the terpolymerization; or

continuously supplying the catalyst and the cocatalyst to the reaction zone during the polymerization with the formation of the activated complex during the terpolymerization; or

any other method suitabla for athylena polymarization can be usad.

[0086] The athylene metallocene copolymerization process according to this invention may thus involve bringing into contact in the reaction zone at least othylene, and a further alpha olefin with the proviso that the further alpha olefin has a total number of carbon higher than 5.

[0087] The ethylene metallocene terpolymerization process according to this invention may thus involve bringing into contact in the reaction zona at least ethylene, a second alpha olefin and a further alpha olefin with the proviso that at least the second alpha olefin has a total number of carbon higher than 5.

[0088] The molecular weight distribution of such terpolymers can vary according to a particular metallocene catalyst employed, a particular co-catalyst employed and a particular mixture of alpha olefins employed.

[0089] The invention will now be described in more detail with reference to the following non limiting examples [0090] In the examples:

(i) Melting behavior, is the Tm meiting temperature and the heat of fusion, wara dataminad on a Parkin Elmar DSC7 fittad with a TAC7/PC instrument controller. The samples were heated from 50°C to 200°C at 20°C/min, held at 200°C for 1 min, cooled to 50°C at a rate of 20°C/min, and held at 50°C for 1 minute after which the maiting curve was recorded between 50°C and 200°C at a heating rate of 10°C/min.

(ii) Molecular waight was determined on a Waters 150 CV GP chromatograph equipped with a data module and computer acquisition system. Determinations were done on polymer samples dissolved at 150°C in 1,24 trichiorobenzañe. Each tray of samples included a polystyrene standard and a NBS 1475a standard in order to check the validity of data against the calibration curve data. Differential Refractive Index was used for detection. These tests were used to determine polydispersity and the molecular weight average viscosity weight.

(iii) Rheological analysis were performed on a Physica MCR-500 rheometer with a temperature control unit. Frequency sweep experiments (oscillatory test) were performed with a parallel plate measuring system (25 mm diameter), under controlled strain conditions, at a constant temperature. The measuring conditions were as follows:

The measuring temperatures wera T = 150, 160, 170 °C.

The strain = 1 %

The rheological values were determined by varying the angular frequency logarithmically from 0.01 to 500\(\ell\)/s The gap between the parallel plates was set to 1 mm.

Measurements were performed on heat stabilised injection molded disks (25 diameter x 2 mm thick) of the polymer.

EXAMPLE 1

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[00st] To a 1-liter stainless steel automated autociave fitted with heating, cooling and stirring facilities, and thoroughly purged with nitrogen, was added 350g polymerization grade cyclohexane and the temperature set at 80°C. To this was added 10 ml of a 30% solution of MAO in tolure presented with 0.1 mg bis n-buryl cyclopentatione zirconium dichloride. Ethylane and a 31′ mass mixture of Fischer-Tropsch derived 1-pentane and 1-butona ware introduced simultaneously at 10m vatas of 25mml. After 50 ml in the monomer flows were stopped, and the reaction continued for another 10 minutes after which the reaction mixture was cooled down. The catalyst was then deactivated by the introduction of 100 ml iso propanel and the surry filtered, washed repeatedly with acctione and dried under vacuum at 70°C. The yield of terpolymer was 80g. The density of the copolymer as measured according to ASTIM D 1506 was 0,9158 g/cm³ and MFI measured according to ASTIM D 1508 was 0,1 dg/min. A GPC measurement was carried out and the M_m, M_m, M_m and polydispensity (P) of the copolymer were 261330, 538870, 395922 and 2,06 raspectivaly, DSC gave a melting temperature of 119 dcs. and a fusion entialsy of 132,0/s. Rhookological determinations were done and the dynamic zero

shear viscosity and the cross-over frequency ω_c were respectively 3.71E5 Pa.s and 4 rad/s at 150 °C; were respectively 3.32E5 Pa.s and 5 rad/s at 160 °C; and were respectively 3.18E5 Pa.s and 5.1 rad/s at 170 °C.

EXAMPLE 2

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[0092] To a 10-fiter stainless steel automated autoclave fitted with heating, cooling and stirring facilities, and thoroughly purged with nitrogen, was added 1500g polymerization grade toluene and the temperature set at 85°C. To this was added 10 mil of a 50% solution of MAD in toluene prereacted with 0.58 mg bis 1-ethyl indenyl zirconium dichloride and 100 mg hydrogen. Ethylene and Fischer-Tropsch derived 1-pentene were introduced simultaneously at flow rates of 5g/min. After 80 min. the monomer flows were stopped and the reaction mixture cooled down over a period of 5 minutes. The catalyst was then deactivated by the Introduction of 500 mil so propanol and the slurry filtered, washed repeatedly with acetone and dried under vacuum at 70°C. The yleid of copolymer containing 2.8% 1-pentene was 490g. The density of the copolymer as measured according to ASTM D 1508 was 0.9350g/cm³ and MFI measured according to ASTM D 1238 was 1.8 domin.

EXAMPLE 3

[0093] To a 10-liter stainless steel automated autoclave fitted with heating, cooling and stirring facilities, and thoroughly purged with hirtogen, was added 2009 optymerization grade toluene and the temperature set at 85°C. In this was added 100 ml of a 30% solution of MAO in toluene prereated with 1.6 mg bis n-buty cyclopentadiene zirconium chloridies. Ethickness and Ethickness Tops and the rest of the set into the set of 12.5 g/s min. After 80 min. the monomer flows were stopped and the reaction mixture cooled down over a period of 60 minutes. The catalyst was then deactivated by the introduction of 500 mll is propanal can the surry filtered, washed repeated with a catone and dried under vacuum at 70°C. The yield of copolymer containing 2% 1-pentiene was 640g. The density of the copolymer as measured according to ASTM to 1565 was 9.803 (g/cm² and MF imeasured according to ASTM to 1565 was 9.803 (g/cm² and MF imeasured according to ASTM to 1565 was 0.803 (g/cm² and MF imeasured according to ASTM to 1565 was needing to 1900 and Firm according to ASTM to 1565 was melting temperature of 115 deep, and a fusion entity of 94 U/g. Rheological determinations were done and the dynamic zero shear viscosity η* was 809 Pa.s at 150 °C, 49 Pa.s at 160 °C and 544 Pa.s at 170 °C.

EXAMPLE 4

[0094] To a 10-liter stainless steel automated autoclave fitted with healing, cooling and stirring facilities, and thooughly purged with hilrogen, was added 2000 polymerazion grade toluene and the temperature set at 85°C. To his was added 100 mi of a 30% solution of MAO in toluene prereacted with 0.8 mg bis n-burly cyclopentacliene zirconium dichloride. Ethieven and Fischer-Tropsch derived 1-pentene were introduced simultaneously at flow rates of 9 gmin and 2.5 g/min respectively. After 80 min. the monomer flows were stopped and the reaction mixture cooled down over a period of 80 minutes. The catalyst was then deactivated by the introduction of 500 mi los propanol and the surry filtered, washed repeatedly with acetone and dried under vacuum at 70°C. The yield of copplymer containing 0.44% 1-pentene was 830g. The density of the copplymer as measured according to ASTM D 1505 was 0.555 gcm³ and MFI measured according to ASTM D 1505 were 2005. Signal of MFI measured according to ASTM D 1505 was 0.555 gcm³ and MFI measured according to the copplymer were \$2045, 211903, 45898 and 2.58 respectively. DSC gave a method to the M_M, M_m, and polydispensity (P) of the copplymer were \$2045, 211903, 45898 and 2.58 respectively. DSC gave a method to the department of 125 deg. and a fusion enthalpy of 143. Jg. Rheological determinations were done and the dynamic zero shear viscosity m², the cross-cover modulus of, the cross-cover frequency ω_c were respectively 1515 Pa.s., 19445 Pa.s., 19445

EXAMPLE 5

[0095] To a 1-liter stainless steel automated autoclave fitted with heating, cooling and stirring facilities, and thoroughly purged with hitrogen, was acided 350g polymer/ation grade to bluene and the temperature set at 80°C. To this was added 1.5 ml of a 30% solution of MAO in tolurene followed 5 minutes later by 2mil of a pre-contacted solution containing 1 ml of a 5°d M solution of a bis-inderly dimethyl silyl zicconium dichloride and 1 ml of a 30% solution of MAO, both in tolurene. Ethylene and Fischer-Tropsch derived 1-hexene were introduced simultaneously at flow rates of 4 ghmin, and 2 ghmin, respectively. After 100 g of ethylene and 50 g of Fisher-Tropsch derived 1-hexene were introduced the monemer flows were stopped and the reaction continued for another 35 minutes. After this period the catalyst was decivited by the introduction of 100 ml iso propanol and the slurny cooled to room temperature under stirring. This slurny was filtered, washed repeatedly with acction and dried under vacuum at 80°C. The yield of copylmer was 125g. The

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density of the copolymer as measured according to ASTM D 1505 was 0.94 g/cm² and MFI measured according to ASTM D 1208 was 0.04 g/cm². As of the copolymer were 96750, 227492, 435490 and 2.35 respectively, DSC gave a melting temperature of 123 deg. and a fusion enthalpy of 112 g/s. Reboolgical eleminations ever done and the dynamic zors observatively 150 and 170 "C and the cross-over complex shear viscosity n° was 1221300 and 1052800 Pas as trespectively 150 and 170 "C and the cross-over complex shear viscosity n° ewas 934820, 750350 and 444480 Pas as trespectively 160, 160 and 170 "C.)

EXAMPLE 6

[0098] To a 10-liter stainless steel automated autoclave fitted with heating, cooling and stirring facilities, and thoroughly purged with nitrogen, was added 2000g polymerization grade toluene and the temperature set at 85°C. To this was added 100 ml of a 30% solution of MAO in toluene followed 5 minutes later by 0.8 mg of a n-butyl cyclopentadiene zirconium dichloride. Ethylene and Fischer-Tropsch derived 1-hexene were introduced simultaneously at flow rates of 9 g/mln, and 2.5 g/mln, respectively. After 720 g of ethylene and 200 g of Fisher-Tropsch derived 1-hexene were introduced the monomer flows were stopped and the reaction continued for another 40 minutes. After this period the catalyst was deactivated by the introduction of 100 ml iso propanol and the slurry cooled to room temperature under stirring. This sturry was filtered, washed repeatedly with acetone and dried under vacuum at 80°C. The yield of copolymer with a composition 0.2%, 1-hexene was 580g. The density of the copolymer as measured according to ASTM D 1505 was 0.931 g/cm3 and MFI measured according to ASTM D 1238 was 0.73 dg/min. A GPC measurement was carried out and the M_n, M_w, M_z and polydispersity (P) of the copolymer were 95376, 243707, 501000 and 2.56 respectively. DSC gave a melting temperature of 126 deg. and a fusion enthalpy of 148 J/g. Rheological determinations were done and the dynamic zero shear viscosity η° , the cross-over modulus G_{n} , the cross-over frequency ω_{n} , the crossover complex shear viscosity n a were respectively 99690 Pa.s, 202840 Pa, 39 rad/s and 7346 Pa.s at 150 °C; were respectively 80554 Pa.s, 198400 Pa, 43 rad/s and 6516 Pa.s at 160 °C; and were respectively 41017 Pa.s. 186260 25 Pa, 52 rad/s and 5079 Pa,s at 170 °C.

EXAMPLE 7

[0097] To a 10-lifer stainless steel automated autoclave fitted with heating, cooling and stirring facilities, and thoroughly purped with introgen, was added 2000 polymerazionic grade toluene and the temperature set at 85°C. To this was added 100 ml of a 10% solution of MAO in toluene prereated with 0.8 mg bis n-buyl cyclopentadiene zirconium dichlorids. Ethylene and a 11′ mass mixture of Fischer-Tropsech derived 1-pentene and 1-hexene were introduced simultaneously at flow rates of 15 g/min. and 3 g/min. respectively. After 50 min. the monomer flows were stopped, and the reaction continued for another 30 minutes. The datalyst was then describated by the introduction of 500 mil as propanol and the situry filtered, washed repeatedly with actorione and dried under vacuum at 70°C. The yield of terpolymer containing 0.17% 1-pentene and 0.23% 1-hexene respectively, was 339g. The density of the copolymer as measured according to ASTM 0.1506 was 0.9497 g/cm² and MR1 measured according to ASTM 0.1288 was 1 dgmin. A GPC measurement was carried out and the M_m, M_m, M_m and polydisperaity (P) of the copolymer were 123059, 28387, 5630328 and 2.31 respectively. DSC gave a melting temperature of 126 deg, and a fusion enthalby of 145 U.g he-cross-over modulus G_c, the cross-over modulus G_c, the cross-over modulus G_c, the cross-over complex shear viscosity n^{*}, eme respectively 4447 Pa. s, 17899 Pa. 3, 2 Trad/s and 37202 Pa. s at 150°C; were respectively 64802 Pa. s, 188640 Pa. 36 rad/s and 7268 Pa.s at 160°C; and were respectively 59435 Pa. s. 22780P Pa. 4170°C.

45 EXAMPLE 8

[0098] To a 10-liter stainless steal automated autoclave fitted with heating, cooling and stirring facilities, and thoroughly purged with nitrogen, was added 2000g polymerization grade toluene and the temperature set at 85°C. To this was added 100 ml of a 10% solution of MAO in toluene prereacted with 0.8 mg bis n-buyly cyclopentacismo zirconium of the object of the control of t

EXAMPLE 9

[0099] -To a 10-liter stainless steel automated autodave fitted with heating, cooling and stirring facilities, and thocoughly purged with introgen, was added 2009 polymerization grade toluene and the temperature set at 85°C. bits was added 100 ml of a 10% solution of MAO in toluene premacted with 0.8 mg bis n-butyl cyclopentadiene zirconium
dichloride. Ethylene and a 1/2 mass mixture of Fischer-Tropsch derived 1-pentene and 1-hexane were introduced
simultaneously at flow rates of 15 g/min. and 5 g/min. respectively. After 50 min. the monomer flows were stopped,
and the reaction continued for another 30 minutes. The catalyst was then deactivated by the introduction of 500 ml is propanol and the sturry filtered, washed repeatedly with acetone and dried under vacuum at 70°C. The yield of they
propanol and the sturry filtered, washed repeatedly with acetone and dried under vacuum at 70°C. The yield of they
under containing 0.15% 1-pentene and 0.23% 1-hexane respectively, was 850g. The density of the copolymer as measured according to ASTM 0.15% was 0.9537 g/cm² and MFI measured according to ASTM 0.15% was 0.9537 g/cm² and MFI measured according to ASTM 0.15% was 0.9537 g/cm² and MFI measured according to ASTM 0.15% was 0.9537 g/cm² and MFI measured according to ASTM 0.15% was 0.9537 g/cm² and MFI measured according to ASTM 0.15% was 0.9537 g/cm² and MFI measured according to ASTM 0.15% was 0.9537 g/cm² and MFI measured according to ASTM 0.15% was 0.9537 g/cm² and MFI measured according to ASTM 0.15% was 0.9537 g/cm² and MFI measured according to ASTM 0.15% was 0.9537 g/cm² and MFI measured according to ASTM 0.15% was 0.9537 g/cm² and MFI measured according to ASTM 0.15% was 0.9537 g/cm² and MFI measured according to ASTM 0.15% was 0.9537 g/cm² and MFI measured according to ASTM 0.15% was 0.9537 g/cm² and MFI measured according to ASTM 0.15% was 0.9537 g/cm² and MFI measured according to ASTM 0.15% was 0.9537 g/cm² and MFI measured according to ASTM 0.15% was 0.9537 g/cm² and MFI measured according to ASTM 0.15% w

5 EXAMPLE 10

[0100] To a 10-liter stainless steel automated autocave fitted with healing, cooling and stirring facilities, and thoroughly purped with nitrogen, was added 2000 polymerization grade toluene and the temperature set at 85°C. To this was added 100 ml of a 10% solution of MAO in toluene prereasted with 0.8 mg bis n-butyl cyclopentadiene zirconium dichloride. Ethylene and a 23 mass mixture of Fischer-Tropsoh derived 1-pentene and 1-hexene were introduced simultaneously at flow rates of 24 grims, and 10 grim, respectively. After 50 min. the monomer flows were stopped, and the reaction continued for another 30 minutes. The catalyst was then deactivated by the introduction of 500 ml lso-propanol and the sturry filtered, washed repeatedly with acetone and dried under vacuum at 70°C. The yield of the proprogramer containing 0.35% 1-hexene respectively, was 780g. The density of the copolymer as measured according to ASTM 0 150% was 0.395 (grow? and MF1 measured according to ASTM 0 1528 was 2.9 dg/m². A GPC measurement was carried out and the M_m, M_m, M_m and polydispersity (?) of the copolymer were 7337, 173645, 354722 and 2.37 respectively, 05C gave a melting temperature of 121 deg. and a fusion entitlept of 127 Ug.

EXAMPLE 11

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[0101] To a 10-liter stainless steel automated autoclave fitted with heating, cooling and stirring facilities, and thorough purged with hitrogen, was added 2009 polymerization grade toluene and the temperature set at 86°C. To this was added 150 ml of a 10% solution of MAO in toluene presented with 0.8 mg bis n-butyl cyclopentacline zirconium dichloride. Ethylene and a 1/1 mass mixture of Fischer-Tropsch derived 1-pentene and 1-hexene were introduced simultaneously at flow rates of 18 g/min. and 12 g/min. respectively. After 50 min. the monomer flows were stopped, and the reaction continued for another 30 minutes. The catalyst was then deactivated by the hitroduction of 500 ml its propanol and the surry filtersch washed repetiedly with acctione and dried under vacuum at 70°C. The yield of terpolymer containing 0.3% 1-pentene and 0.35% 1-hexene respectively, was 810g. The density of the copolymer as measured according to ASTM D 1508 was 0.389 g/m² and MFI measured according to ASTM D 1508 was 0.389 g/m² and MFI measured according to ASTM D 1508 was 1.2 g/m² and MFI measured according to ASTM D 1508 was 1.2 g/m² and MFI measured according to ASTM D 1508 was 1.2 g/m² and MFI measured according to ASTM D 1508 was 1.2 g/m² and MFI measured according to ASTM D 1508 was 1.2 g/m² and MFI measured according to ASTM D 1508 was 1.2 g/m² and MFI measured according to ASTM D 1508 was 1.2 g/m² and MFI measured according to ASTM D 1508 was 1.2 g/m² and MFI measured according to ASTM D 1508 was 1.2 g/m² and MFI measured according to ASTM D 1508 was 1.2 g/m² and MFI measured according to ASTM D 1508 was 1.2 g/m² and MFI measured according to ASTM D 1508 was 1.2 g/m² and MFI measured according to ASTM D 1508 was 1.2 g/m² and MFI measured according to ASTM D 1508 was 1.2 g/m² and MFI measured according to ASTM D 1508 was 1.2 g/m² and MFI measured according to ASTM D 1508 was 1.2 g/m² and MFI measured according to ASTM D 1508 was 1.2 g/m² and MFI measured according to ASTM D 1508 was 1.2 g/m² and MFI measured according to ASTM D 15

EXAMPLE 12

45 [0102] To a 1-lier stainless steel automated autoclave fitted with heating, cooling and strring facilities, and thoroughly purged with throtogen, was added 350g polymerization grade toluene and the temperature set at 80°C. To this was added 1.5 ml of a 30% solution of MAO in toluene followed 5 minutes later by 2m of a precontacted solution containing 1 ml of a 5% solution of a bis-inderyl dimethylayly zirconium dichloride and 1 ml of a 30% solution of MAO, both in Ollone. Ethylene and Fischer-Tropsech derived 1-heptene were introduced simultaneously at flow rates of 4 g/min. and 92 g/min. respectively. After 100 g of ethylene and 50 g of Fisher-Tropsech derived 1-heptene were introduced simultaneously at flow rates of 4 g/min. and 92 g/min. respectively. After 100 g of ethylene and 50 g of Fisher-Tropsech derived 1-heptene were introduced to monomer flows were stopped and the reaction continued for another 35 minutes. After this period the catalyst was deactivated by the introduction of 100 ml is porpopanol and the subrry cocked for room temperature under stirring. This slurry was filtered, washed repeatedly with acetione and dried under vacuum at 80°C. The yield of copolymer was 95g. The density of the copolymer as measured according to ASTM D 1505 was 0.956 g/m²n and MFI measured according of the copolymer were 785s. (170303, 32285g and 2.23 respectively. DSC gave a melting temperature of 126 deg. and a fusion enthalby of 55 u/g. Rheological determinations were done and the dynamic zero shear viscosity n² and the cross-over complex shear viscosity n² a

and 5312 Pa.s at 160 °C; and were respectively 80670 and 4350 Pa.s at 170 °C.

EXAMPLE 13

5 [0103] To a 1-liter stainless steel automated autoclave fitted with healing, cooling and stirring facilities, and thoroughly purgod with Introgen, was added 350g polymerization grade toluren and the temperature set at 80°C. To this was added 1.5 ml of a 30% solution of MAO. In toluren's clowed 5 minutes later by 2ml of a pre-contacted solution containing 1 ml of a 5°3 M solution of a bis-indenyl dimethyl silyl zircontium dichloride and 1 ml of a 90% solution of MAO. both in tolurene. Ethylene and a 1/1 mixture of Fisher-Tropsch derived 1-heptiene and 1-pentiene were introduced alminated ously at flow rates of 4 g/min. and 2 g/min. respectively, After 100 g of ethylene and 50 g of the mixture of Fisher-Tropsch derived 1-heptiene and the reaction continued for another 35 minutes. After this period the catalyst was describated by the introduction of 100 ml iso propanol and the submy cooled to room temperature under stirring. This surry was filtered, washed repeatedly with accrotion and dried under vacuum at 80°C. The yield of terpolymer was 82g. The density of the copolymer as measured according to ASTM D 1250 was 0.5 tig/min. A GPC measurement was carried out and the M_{in}, M_{in}, M_{in} and polydisporatity (P) of the copolymer vere 48305, 131286, 311478 and 2.72 respectively. DSC gave a metting temperature under and fusion enthalpy of 24 J/g.

EXAMPLE 14

[0104] To a 1-liter stainless steel automated autoclave fitted with heating, cooling and stirring facilities, and thoroughly purged with introgen, was added 350g polymorization grade to bluene and the temperature set at 80°C. To those was added 1.5 ml of a 30% solution of MAO in toluren followed 5 minutes later by 2ml of a precontacted solution containing 1 ml of a 5³ M solution of a bis-indenyl dimethylally zirconium dichiordic and 1 ml of a 50% solution of MAO, both in toluren. Ethylane and Flischer-Tropsch derived 1-octene were introduced simultaneously at 10 was trase both of 4 g/min. After 100 g of ethylene and 100 g of Fisher-Tropsch derived 1-octene were introduced, the monomer flows were stopped and the reaction continued for another 35 minutes. After this period the catalyst was deactivated by the Introduced of 10 ml iso propanol and the sturry cooled to room temperature under stirring. This sturry was filtered, washed repeatedly with acctone and cride under vacuum at 80°C. The yield of copolymer, containing 9.8% 1-octene, was 170%. The density of the copolymer as measured according to ASTM D 1505 was 0.88 g/cm³ and MFI measured according to ASTM D 1505 was 0.88 g/cm³ and MFI measured according to ASTM D 1505 was 0.88 g/cm³ and MFI measured according to ASTM D 1505 was 0.88 g/cm³ and MFI measured according to ASTM D 1505 was 0.88 g/cm³ and MFI measured according to ASTM D 1505 was 0.88 g/cm³ and MFI measured according to ASTM D 1505 was 0.88 g/cm³ and MFI measured according to ASTM D 1505 was 0.88 g/cm³ and MFI measured according to ASTM D 1505 was 0.88 g/cm³ and MFI measured according to ASTM D 1505 was 0.88 g/cm³ and MFI measured according to ASTM D 1505 was 0.88 g/cm³ and MFI measured according to ASTM D 1505 was 0.88 g/cm³ and MFI measured according to ASTM D 1505 was 0.88 g/cm³ and MFI measured according to ASTM D 1505 was 0.88 g/cm³ and MFI measured according to ASTM D 1505 was 0.88 g/cm³ and MFI measured according to ASTM D 1505 was 0.88 g/cm³ and MFI measured according to ASTM D 1505 was 0.88

35 EXAMPLE 15

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[0105] To a 1-lier stainless ateal automated autoclave fitted with heating, cooling and stirring facilities, and thoroughly purged with introgen, was added 350g polymerization grade to bluene and the temperature set at 80°C. To this was added 1.5 ml of a 30% solution of MAO in toluren followed 5 minutes later by 2ml of a precontacted solution containing 1 ml of a 5°3 M solution of a bis-inderly dimethyshigh zitconium dichloride and 1 ml of a 30% solution of MAO, both in toluren. Ethylene and Fischer-Tropsch derived 1-octene were introduced simultaneously at thow trates both of 4 g/ml. After 100 g of shrylene and 100 g of Fisher-Tropsch derived 1-octene were introduced the monomer flows were stopped and the reaction continued for another 35 minutes. After this period the catalyst was deactivated by the introduction of 100 ml iso propanol and the siture occoled to room temperature under stirring. This situry was filtered, washed repeatedly with acctone and dried under vacuum at 80°C. The yield of copolymer, containing 13.4% 1-octene, was 1505. The density of the copolymer as measured according to ASTM D 1505 was 0.84 g/cm³ and MFI measured according to ASTM D 1505 was 0.84 g/cm³ and MFI measured according to ASTM D 1505 was 0.84 g/cm³ and MFI measured according to ASTM D 1505 was 0.84 g/cm³ and MFI measured according to ASTM D 1505 was 0.84 g/cm³ and MFI measured according to ASTM D 1505 was 0.84 g/cm³ and MFI measured according to ASTM D 1505 was 0.84 g/cm³ and MFI measured according to ASTM D 1505 was 0.84 g/cm³ and MFI measured according to ASTM D 1505 was 0.84 g/cm³ and MFI measured according to ASTM D 1505 was 0.84 g/cm³ and MFI measured according to ASTM D 1505 was 0.84 g/cm³ and MFI measured according to ASTM D 1505 was 0.84 g/cm³ and MFI measured according to ASTM D 1505 was 0.84 g/cm³ and MFI measured according to ASTM D 1505 was 0.84 g/cm³ and MFI measured according to ASTM D 1505 was 0.84 g/cm³ and MFI measured according to ASTM D 1505 was 0.84 g/cm³ and MFI measured according to ASTM D 1505 was 0.84 g

EXAMPLE 16

[D168] To a 1-liter stainless steel automated autoclave fitted with heating, cooling and stirring facilities, and throughly purged with introgen, was added 350g polymerization grade cyclohexane and the temperature set at 80°C. To this was added 10 mil of a 30% solution of MAO in toluren prereacted with 0.1 mg bis n-buyly cyclopentadiene zirconium dichloride. Ethylene and a 1.2 mass mixture of Fischer-Tropsch derived 1-pention and 1-octane were introduced similar neously at flow rates of 2 g/min. and 1.5 g/min. respectively. After 50 min. the monomer flows were stopped, and the neaction continued for another 10 minutes after which the reaction mixture was cooled down. The catalyst was the

deactivated by the Introduction of 100 miles propanol and the stury filtered, washed repeatedly with section and dried under vacuum at 70°C. The yield of timpolymer, containing 1.0°4, 1 penteins and 0.4% 1-oction, vao 70g. The deneity.

of the copolymer as measured according to ASTM 0.1505 was 0.9191 g/cm³ and MFI measured according to ASTM 0.1288 was 750/min. A QFC measurement was carried out and the M_N, M_c, M_c and polysipsensity (Po 1 the copolymer were 52789, 147903, 307128 and 2.0 respectively, Pheological determinations were done according to description and the Carreau-Galintienre qualition parameters m₀, a, b, pewer respectively 6.765-F2 ns., 45E-4, 0.38, 4.1, at 150°C; were respectively 6.04E-5 Pa.s., 9.88E-4, 0.39, 3.2, at 160°C; and were respectively 5.31E-5 Pa.s., 1.33E-3, 0.42.2.9, at 170°C

10 EXAMPLE 17

[0107] To a 1-liter stainless steel automated autociave fitted with heating, cooling and stirring facilities, and thoroughly purged with introgen, was added 350 polymerization grade to toutnee and the temperature set at 80°C. To this was added 1.5 ml of a 30% solution of MAO in toluene followed 5 minutes later by 2ml of a pre-contacted solution containing 1 ml of a 5°d is obtition of a bis-inderly dimethy silly zilconium dichioride and 1 ml of a 30% solution of MAO, both in toluene. Ethylene and Fischer-Tropach derived 1-nonene were introduced simultaneously at flow rates of 4gmin, and 2 gmin, respectively. After 100 g of ethylene and 50 g of Fisher-Tropach derived 1-nonene were introduced the moner flows were stopped and the reaction continued for another 35 minutes. After this period the catalyst was described by the introduction of 100 ml iso propanol and the sturry cooled to room temperature under stirring. This sturry was filtered, washed repeatedly with acetone and dried under vacuum at 80°C. The yield of copolymer was 1146, The density of the copolymer as measured according to ASTM D 1505 was 0.90 g/cm² and MFI measured according to ASTM D 1505 was 0.90 g/cm² and MFI measured according to ASTM D 1505 was 0.90 g/cm² and MFI measured according to the copolymer were 101377, 252832, 446233 and 2.49 respectively. DSC gave melting temperature of 128 deg. and a fusion enthalpy of 157 J/g. Rheological acterimations were done and the Carraau-Galheliner equation parameters $1_{\rm Hz}$ 8, b, p and b/p were respectively 1.18E-4 Pas, 9.44), 3.41, 0.21, 1.51, 0.81 150 °C; over respectively 1.18E-4 Pas, 9.49, 1,3.41, 0.21, 1.51, 0.81 150 °C; were respectively 1.18E-4 Pas, 9.49, 1.24, 0.24, 8.34, 8.41 the 100 °C; and were respectively 1.18E-4 Pas, 9.49, 1.24, 0.24, 8.34, 8.41 the 100 °C; and were respectively 1.18E-4 Pas, 9.49, 1.24, 0.24, 8.34, 8.41 the 100 °C; and were respectively 1.24 constant of the proper temperature of 128 deg. and 100 °C; and were respectively 1.24 constant of the proper temperature of 128 deg. a

EXAMPLE 18

[0108] To a 1-liter stainless steel automated autociave fitted with heating, cooling and stirring facilities, and thoroughly purged with nitrogen, was added 150g polymerization grade toluene and the temperature set at 80°C. To this was added 1.5 m of a 30% solution of MAO in toluene followed 5 minutes later by 2m of a pre-constacted solution containing 1 mil of a 5° M solution of a bis-indenyl dimethyl sily zirconium dichloride and 1 mil of a 30% solution of MAO, both in toluene. Eithylene and a 50°G mixture of Fisher-Tropsch derived 1-nonene and 1-pentene were introduced simultanes necessary and 1 mil of 1

45 Claims

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- A polymer of ethylene as a first monomeric component, with a high carbon number linear alpha olefin having at least five carbon atoms as a second monomeric component, with at least one of the monomeric components being Fischer-Tropsch derived so that it includes at least one other olefinic component.
- A polymer according to Claim 1, wherein a plurality of the other cleffnic components are present in the Fischer-Tropsch derived monomeric component, with the molar proportion of other cleffnic components in the Fischer-Tropsch derived monomeric component being from 0,002% to 2%.
- 3. A polymer according to Claim 2, wherein the second monomeric component is Fischer-Tropsch derived; and wherein the high carbon number linear alpha olefin is 1-pentiene, with the other olefinic components constituting about 0,5% of the second monomeric component and comprising

- 2-methyl-1-butene; and/or
- branched elefine having a carbon number of 5; and/or
- internal olefins having a carbon number of 5; and/or cyclic olefins having a carbon number of 5;

and/or

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wherein the high carbon number linear alpha olefin is 1-hexene, with the other olefinic components present in the second monomeric component comprising

- branched olefins having a carbon number of 6; and/or
- internal olefins having a carbon number of 6; and/or
- cyclic olefins having a carbon number of 6;

and/or

15 wherein the high carbon number linear alpha olefin is 1-heptene, with the other olefinic components present in the second monomeric component comprising

- branched olefins having a carbon number of 7; and/or
- internal olefins having a carbon number of 7;

wherein the high carbon number linear alpha olefin is 1-octene, with the other olefinic components present in the second monomeric component comprising

- branched olefins having a carbon number of 8; and/or
- Internal olefins having a carbon number of 8;

wherein the high carbon number linear alpha olefin is 1-nonene, with the other olefinic components present in the second monomeric component comprising

- branched olefins having a carbon number of 9; and/or
- internal olefins having a carbon number of 9.
- 4. A polymer according to Claim 3, wherein the ratio of the molar proportion of ethylene to the molar proportion of 35 the high carbon number linear alpha olefin is from 99,9:0,1 to 80:20.
 - 5. A polymer according to Claim 3 or Claim 4, which has a melt flow rate of from 0.01 to 100g/10mInutes and/or which has a density in the range 0.835 to 0.950g/cc.
 - 6. A polymer according to any one of Claims 3 to 5 inclusive, which has a power exponent b which complies with the following equations:

At 150°C, b ≥ 0.0437[C] + 0.2013

At 160°C, b ≥ 0.0308fCl + 0.2138

At 170°C, b ≥ 0.0308[C] + 0.2538,

where b is a Carreau-Gahleitner parameter, and [C] is the high carbon number linear alpha olefin content in mole percentage; and/or 55

which has a power exponent p which complies with the following equations:

At 150°C, p ≥ -1.2877[C] + 6.8666

At 160°C, p ≥ -1.1233[C] + 6.3942

At 170°C, p ≥ -1.1507[C] + 6.3063.

where p is a Carreau-Gahleitner parameter, and is the high carbon number linear alpha olefin content in mole percentage; and/or

which has a power exponent n which complies with the following equations:

At 150°C, n ≥ 0.2995[C] - 0.8328

At 160°C, n > 0.3011 [C]- 0.8435

At 170°C, n ≥ 0.2942[C] - 0.8115

where n is a Carreau-Gahleitner parameter, and [C] is the high carbon number linear alpha olefin content in mote percentage, and/or which compiles with the following equations:

which complies with the following equations

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At 150°C, 1/MFI ≥ 1.5364e^{-1E-08η°}

At 160°C, 1/MFI ≥ 1.5486e*1E-06η*

At 170°C, 1/MFI ≥ 1.5513e^{-1E-06}η*

where MFI is the melt flow index and η° is the dynamic zero shear viscosity.

- 7. A polymer according to any one of Claims 3 to 5 inclusive, which is that obtained by reacting at least ethylene and the high carbon number Fischer-Tropsch derived linear alpha olefin in one or more reaction zones, while maintaining the reaction zone(s) at a pressure between atmospheric pressure and 5000kg/cm², and at a temperature between ambient and 300°C, in the presence of a metalliceene catalyst, or a catalyst system comprising a metallocene catalyst and a cocatalyst.
- 8. A process for producing a polymer, which comprises reacting at least a first monomeric component comprising ethylene and a second monomeric component comprising a high carbon number linear alpha olelfn having at least five carbon atoms, and wherein at least one of the monomeric components is Fischer-Tropsch derived so that it contains also one or more other olefinic components, in one or more reaction zones, while maintaining the reaction zone(s) at a pressure between atmospheric pressure and 5000kg/cm², and at a temperature between ambient and 500°C, in the presence of a catalyst, or a catalyst system comprising a catalyst and a cocatalyst.
- A process according to Claim 8, wherein the second monomeric component is Fischer-Tropsch derived; and wherein the high carbon number linear alpha clefin is 1-pentene, with the other clefinic components constituting about 0,5% of the second monomeric component and comprising
 - 2-methyl-1-butene; and/or
 - branched olefins having a carbon number of 5; and/or
 - internal olefins having a carbon number of 5; and/or
 - cyclic olefins having a carbon number of 5;

and/or

wherein the high carbon number linear alpha olefin is 1-hexene, with the other olefinic components present in the

second monomeric component comprising

- branched olefins having a carbon number of 6; and/or
 internal olefins having a carbon number of 6; and/or
- cyclic olefins having a carbon number of 6;

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wherein the high carbon number linear alpha olefin is 1-heptene, and wherein the second monomeric component is Fischer-Tropsch derived, with the other clefinic components present in the second monomeric component comorising

- branched olefins having a carbon number of 7: and/or
- internal olefins having a carbon number of 7:

and/or

wherein the high carbon number linear alpha olefin is 1-octene, with the other olefinic components present in the second monomeric component comprising

branched olefins having a carbon number of 8; and/or
 internal olefins having a carbon number of 8;

and/or

wherein the high carbon number linear alpha olefin is 1-nonene, and wherein the second monomeric component is Fischer-Tropsch derived, with the other olefinic components present in the second monomeric component comprising

- branched olefins having a carbon number of 9; and/or
- · internal olefins having a carbon number of 9.
- 30 10. A process according to Claim 8 or Claim 9, wherein the catalyst is a metallocene catalyst.
 - 11. A polymer of ethylene as a first monomeric component, with a first high carbon number linear alpha olefin having at least five carbon atoms as a second monomeric component and with a second different high carbon number linear alpha olefin having at least four carbon atoms as a third monomeric component, with at least one of the monomeric components being Fischer-Tropsch derived so that it contains also at least one other olefinic component.
 - 12. A polymer according to Claim 11, wherein a plurality of the other olefinic components are present in the Fischer-Tropsch derived monomeric component, with the molar proportion of other olefinic components in the Fischer-Tropsch derived monomeric component beling from 0.002% to 2%.
 - 13. A polymer according to Claim 12, wherein the second monomeric component is Fischer-Tropsch derived; and wherein the high carbon number linear alpha olefin of the second monomeric component is 1-pentene, with the other olefinic components constituting about 0.5% of the second monomeric component and com
 - 2-methyl-1-butene; and/or
 - branched olefins having a carbon number of 5; and/or
 - internal olefins having a carbon number of 5; and/or
- cyclic olefins having a carbon number of 5;

and/or

wherein the high carbon number linear alpha olefin of the second monomeric component is 1-hexene, with the other olefinic components present in the second monomeric component comprising

- branched olefins having a carbon number of 6; and/or
 - internal olefins having a carbon number of 6; and/or
 - cyclic olefins having a carbon number of 6:

and/or

wherein the high carbon number linear aipha otefin of the second monomeric component is 1-heptene, with the other otefinic components present in the second monomeric component comprising

- branched olefins having a carbon number of 7; and/or
 - internal olefins having a carbon number of 7:

and/or

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wherein the high carbon number linear alpha olefin of the second monomeric component is 1-octene, with the other olefinic components present in the second monomer component comprising

- branched olefins having a carbon number of 8; and/or
- internal olefins having a carbon number of 8;

and/or

wherein the high carbon number linear alpha olefin of the second monomeric component is 1-nonene, with the other olefinic components present in the second monomeric component comprising

- branched olefins having a carbon number of 9: and/or
- internal olefins having a carbon number of 9.
- 14. A polymer according to Claim 12 or Claim 13, wherein the ratio of the molar proportion of ethylene to the sum of the molar proportions of the high carbon number linear alpha cleffins is from 99,9.0,1 to 80.20 and/or wherein the ratio of the molar proportions of the different high carbon number linear alpha cleffins is from 0,1:99,9 to 99,9.0,1.
- 15. A polymer according to Claim 12, which is a terpolymer of ethylene with 1-butene as the third monomeric component and 1-pentene, 1-hexene, 1-heptene, 1-octene, or 1-nonene as the second monomeric component.
- 16. A polymer according to Claim 12, which is a terpolymer of ethylene with 1-pentene as the third monomeric component and 1-hexene, 1-beptene, 1-octene, or 1-nonene as the second monomeric component.
 - 17. A polymer according to Claim 12, which is a terpolymer of ethylene with 1-hexene as the third monomeric component and 1-heptene, 1-octene, or 1-nonene as the second monomeric component.
- 35 18. A polymer according to Claim 12, which is a terpolymer of ethylene with 1-heptene as the third monomeric component and 1-octene, or 1-nonene as the second monomeric component.
 - 19. A polymer according to Claim 12, which is a terpolymer of ethylene with 1-octene as the third monomeric component and 1-nonene as the second monomeric component.
 - 20. A polymer according to any one of Claims 12 to 19 Inclusive, which has a melt flow rate of from 0.01 to 100g/ 10minutes and/or which has a density in the range 0.835 to 0.950g/cc.
 - 21. A polymer according to any one of Claims 12 to 20 inclusive, which has a power exponent b which complies with the following equations:

At 150°C, b ≥ 0.037[C] + 0.2052

At 160°C, b ≥ 0.0395[C] + 0.2342

At 170°C, b ≥ 0.0494[C] + 0.2202,

where b is a Carreau-Gahleitner parameter, and [C] is the high carbon number linear alpha olefin content in mole percentage; and/or

which has a power exponent p which complies with the following equations:

At 150°C, p ≥ -0.4075[C] + 3.4135

At 160°C, p ≥ -0.5016[C] + 3.732

At 170°C, p ≥ -0.9091 [C] + 5.3455

where p is a Carreau-Gahleitner parameter, and [C] is the high carbon number linear alpha olefin content in mole percentage; and/or

which has a power exponent n which complies with the following equations:

At 150°C, n ≥ 0.169[C] - 0.3439

At 160°C, n ≥ 0.1765[C] - 0.4004

At 170°C, n ≥ 0.242[C] - 0.6489

where n is a Carreau-Gahleitner parameter, and [C] is the high carbon number linear alpha olefin content in mole percentage; and/or which complies with the following equations:

At 150°C, 1/MFI ≥ 0.123e^{2E-05η°}

At 160°C, 1/MFI ≥ 0.1239e^{2E-05η°}

At 170°C, 1/MFI ≥ 0.1275e^{2E-05}η°.

where MFI is the melt flow index and η° is the dynamic zero shear viscosity.

- 22. A polymer according to any one of Claims 11 to 21 inclusive, which is that obtained by reacting at least ethylene, the first high carbon number linear alpha olefin and the different high carbon number linear alpha olefin in one or more reaction zones, while maintaining the reaction zone(s) at a pressure between atmospheric pressure and 5000kg/cm², and at a temperature between ambient and 300°C, in the presence of a catalyst, or a metallocene catalyst system comprising a metallocene catalyst.
- 23. A process for producing a polymor, which comprises reacting at least ethylene as a first monomeric component, a first high carbon number linear alpha oldin having at least five carbon atoms as a second monomeric component and a different high carbon number linear alpha olefin having at least four carbon atoms as third monomeric component, with at least one of the monomeric components being Fischer-Tropsch derived so that it contains an or or more reaction zones, while maintaining the reaction zones (so) at a pressure between atmospheric pressure and 5000kg/cm², and at a temperature between ambient and 300°C, in the pressure of a catalyst, or a catalyst at yetom compfising a catalyst and a cocatalyst.
 - 24. A process according to Claim 23, wherein the second monomeric component is Fischer-Tropsch derived; and wherein the first high carbon number linear alpha lollin is 1-pentene, with the other olefinic components constituting about 0,5% of the second monomeric component and comprising
 - 2-methyl-1-butene; and/or

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- branched olefins having a carbon number of 5; and/or
- internal olefins having a carbon number of 5; and/or

- cyclic olefins having a carbon number of 5;

and/or

- wherein the first high carbon number linear alpha olefin is 1-hexene, with the other olefinic components in the second monomeric component comprising
 - branched olefins having a carbon number of 6; and/or
 internal olefins having a carbon number of 6; and/or
 - cyclic olefins having a carbon number of 6;

and/or

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wherein the first high carbon number linear alpha olefin is 1-heptene, with the other olefinic components in the second monomeric component comprising

- branched olefins having a carbon number of 7; and/or
- internal olefins having a carbon number of 7;

and/or

wherein the first high carbon number linear alpha olefin is 1-octene, with the other olefinic components in the second monomeric component comprising

- branched olefins having a carbon number of 8; and/or
- Internal olefins having a carbon number of 8;

25 and

wherein the first high carbon number linear alpha olefin is 1-nonene, with the other olefinic components in the second monomeric component comprising

- branched olefins having a carbon number of 9; and/or
 - internal olefins having a carbon number of 9.

25. A process according to Claim 23 or Claim 24, wherein the catalyst is a metallocene catalyst.



European Patent Office

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